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(71) Applicant:
Agouron Pharmaceuticals, Inc.
La Jolia, CA 92037 (US)

(72) Inventors:

 Chen, Ping San Diego, California 92129 (US)

 Kan, Chen-Chen, Keck Graduate Inst. of A.L.S. Claremont, California 91711 (US)

 Luo, Chun Irvine, California 92206 (US)

 Margosiak, Stephen Escondido, California 92025 (US)

O'Connor, Patrick
 San Diego, California 92130 (US)

Tempczyk-Russel, Anna
 San Diego, California 92130 (US)

Nguyen, Binh
 San Diego, California 92130 (US)

Sarup, Jay Chand
 San Diego, California 92122 (US)

Gaur, Smita
 San Diego, California 92129 (US)

Anderson, Mark Brian
 Orinda, California 94563 (US)

Deng, Ya-Li
 San Diego, California 92130 (US)

Lundgren, Karen
 San Diego, California 92109 (US)

Register, James
 San Diego, California 92192 (US)

(74) Representative:
Hofmann, Harald et al
Sonnenberg Fortmann,
Patent- und Rechtsanwälte,
Herzopgspitalstrasse 10
80331 München (DE)

Remarks:

A request for correction of the description has been filed pursuant to Rule 88 EPC. A decision on the request will be taken during the proceedings before the Examining Division (Guidelines for Examination in the EPO, A-V, 3.).

(54) Catalytic domain of the human effector cell cycle checkpoint protein kinase, Chk1, materials and methods for identification of inhibitors thereof

(57) The present invention relates to the identification, isolation and purification of the catalytic domain of the human effector checkpoint protein kinase (hChk1). A 1.7 crystal structure of the hChk1 kinase domain in the active conformation is reported herein. The kinase domain of hChk1 and its associated crystal structure is described for use in the discovery, identification and characterization of inhibitors of hChk1. This structure provides a three-dimensional description of the binding site of the hChk1 for structure-based design of small molecule inhibitors thereof as therapeutic agents. Inhibitors of hChk1 find utility in the treatment of hyperproliferative disorders such as HIV and cancer.

[0001] This application claims priority from co-pending United States Provisional Application Serial Number 60/162,887, filed November 1, 1999, the contents of which are incorporated by reference herein in their entirety.

FIELD OF THE INVENTION

[0002] The present invention generally relates to cell cycle checkpoint kinases which are essential to cellular DNA damage responses and coordinating cell cycle arrest. The checkpoint kinases play a role in the surveillance and response to DNA damage. The damage may result from external or internal forces. Such forces include but are not limited to errors in replication, DNA base damage, DNA strand breaks, or exposure to radiation or cytotoxic chemicals. These checkpoint kinases are integral in the regulatory pathways leading to cell cycle arrest and apoptosis following DNA damage, giving the cell notice and time to correct lesions prior to the initiation of replication and chromosome separation. The present invention more specifically relates to the isolation and purification of the catalytic domain of the human effector checkpoint protein kinase (hChk1) and its use in the discovery, identification and characterization of inhibitors of same.

BACKGROUND

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[0003] Cell growth, division and death is essential to the life cycle of multi-celled organisms. These processes and their regulation are strikingly similar across all eukaryotic species. Somatic cell division consists of two sequential processes: DNA replication followed by chromosomal separation. The cell spends most of its time preparing for these events in a growth cycle (interphase) which in turn consists of three subphases: initial gap (G_1) , synthesis (S), and secondary gap (G_2) . In G_1 , the cell, whose biosynthetic pathways were slowed during mitosis, resumes a high rate of biosynthesis. The S phase begins when DNA synthesis starts and ends when the DNA content of the nucleus has doubled. The cell then enters G_2 , which lasts until the cell enters the final phase of division, mitotic (M). The M phase begins with nuclear envelope breakdown, chromosome condensation and formation of two identical sets of chromosomes which are separated into two new nuclei. This is followed by cell division (cytokinesis) in which each nuclei is separated into two daughter cells, which terminates the M phase and marks the beginning of interphase for the new cells.

The sequence in which the cell cycle events proceed is tightly regulated such that the initiation of one cell cycle event is dependent upon the successful completion of the prior cell cycle event. The process of monitoring genome integrity and preventing cell cycle progress in the event of DNA damage has been described as a 'cell cycle checkpoint' (Hartwell, LH et at., *Science*, 246:629-634 (1989); Weinert et al., *Genes and Dev.*, 8:652 (1994)]. Cell cycle checkpoints consist of signal transduction cascades which couple DNA damage detection to cell cycle progression. Checkpoints are control systems that coordinate cell cycle progression by influencing the formation, activation and subsequent inactivation of the cyclin-dependent kinases. Checkpoint enzymes are responsible for maintaining the order and fidelity of events of the cell cycle by blocking mitosis in response to unreplicated or damaged DNA. These enzymes prevent cell cycle progression at inappropriate times, maintain the metabolic balance of cells while the cell is arrested and in some instances can induce apoptosis (programmed cell death) when the requirements of the checkpoint havenot been met (O'Connor, PM, *Cancer Surveys*, 29, 151-182 (1997); Nurse, P, *Cell*, 91, 865-867 (1997); Hartwell, LH et al., *Science*, 266, 1821-1828 (1994); Hartwell, LH et al., *Science*, 246, (1989), supra).

[0005] One series of checkpoints monitors the integrity of the genome. Upon sensing DNA damage, these "DNA damage checkpoints" block cell cycle progression in G₁ & G₂ phases, and slow progression through S phase (O'Connor, PM, Cancer Surveys, 29 (1997), supra; Hartwell, LH et at, Science, 266, (1994), supra). This action enables DNA repair to be completed before replication of the genome and subsequent separation of this genetic material into new daughter cell takes place.

[0006] Various mutations associated with malignancy affect the cancer cells ability to regulate checkpoints, allowing cells with DNA damage the increased likelihood to continue replicating and to escape damage-mediated apoptosis. These factors contribute to the genomic instability which drives the genetic evolution of human cancers and contributes to the resistance of cancer cells to most current chemotherapy and radiotherapy intervention.

[0007] Due to abnormalities in the p53 tumor suppressor pathway, most cancer cells lack a functional G_1 checkpoint control system. This makes them particularly vulnerable to abrogation of the last remaining barrier protecting them from the cancer killing effects of DNA damaging agents: the G_2 checkpoint. The G_2 DNA damage checkpoint ensures maintenance of cell viability by delaying progression into mitosis in cells that have suffered genomic damage. The G_2 checkpoint is controlled by cell cycle checkpoint pathways which inhibit mitosis if previous events are incomplete or if the DNA is damaged. This regulation control system has been conserved from yeast to humans. Important in this conserved system is a kinase, Chk1 (or p56Chk1), which transduces signals from the DNA damage sensory complex to inhibit activation of the cyclin B/Cdc2 kinase which promotes mitotic entry (Peng, CY et al, *Science*, 277, 1501-1505

- (1997); Sanchez Y, et al., *Science*, **277**, 1497-1501 (1997); Walworth, N et al., *Nature*, **363**(6427), 368-71 (May 27, 1993); al-Khodairy et al., *Mol Biol Cell*, **5**(2):147-60 (Feb, 1994); Carr et al., *Curr Biol.*, **5**(10): 1179-90 (Oct. 1, 1995)). The repair checkpoint kinase, Chk1, regulates Cdc25, a phosphatase that activates Cdc2. Thus, Chk1 serves as the direct link between the G₂ checkpoint and the negative regulation of Cdc2.
- [0008] Inactivation of Chk1 has been shown to both abrogate G₂ arrest induced by DNA damage inflicted by either anticancer agents or endogenous DNA damage, as well as, result in preferential killing of the resulting checkpoint defective cells (Nurse, P, *Cell*, 91, (1997), <u>supra;</u> Weinert, T, *Science*, 277, 1450-1451 (1997); Walworth, N et al., *Nature*, 363, (1993) <u>supra;</u> al-Khodairy et al., *Molec. Biol. Cell*, 5, (1994), <u>supra;</u> Wan, S et al., *Yeast*, 15(10A), 821-8 (Jul. 1999)).
- [0009] The fact that cancer cells have also been shown to be more vulnerable to G₂ checkpoint abrogation has encouraged the pursuit of G₂ checkpoint abrogating drugs (Wang, Q et al., PNAS 96: 3706-3711 (1999); Fan, S et al., Cancer Res., 55, 1649-1654 (1995); Powell, SN et al., Cancer Res., 55, 1643-1648 (1995); Russell, KJ et al., Cancer Res., 55, 1639-1642 (1995); Wang, Q et al., J Natl Cancer Inst., 88, 956-967 (1996)). Such checkpoint abrogating drugs could improve the killing of tumors exposed to DNA damaging events including that inflicted by therapeutic agents, hypoxic-stress induced because of a limited blood supply (anti-angiogenic agents), or endogenous DNA damage arising as a consequence of a cancer cell's inherent genomic instability. Selective manipulation of checkpoint control in cancer cells can afford broad utilization in cancer chemotherapeutic and radiotherapy regimens and may in addition, offer a common hallmark of human cancer "genomic instability" to be exploited as the selective basis for the destruction cancer cells.
- [0010] A number of lines of evidence place Chk1 as a pivotal target in DNA damage checkpoint control. However, Chk1 is a difficult enzyme to study because the full length protein is not the most active form of Chk1. While others have examined the nucleotide and amino acid sequence of the full-length checkpoint kinase and estimated the location of the kinase domain, there is a need for the isolation and purification of the kinase domain of Chk1 and the maintenance of its catalytically active conformation.

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SUMMARY OF THE INVENTION:

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[0011] The generation, kinetic characterization, and structure determination of the kinase domain of the human Chk1 protein is disclosed herein. The domain begins between residues 1 and 16 and terminates between residues 265 and 291 of the full length protein [SEQ ID NO. 2] which comprises 476 amino acids. The domain preferably extends from residues 1-265, more preferably from residues 1-289.

[0012] The invention relates to an isolated, purified polynucleotide which encodes the active conformation of the human Chk1 kinase or an active kinase analog thereof. The polynucleotide may be natural or recombinant.

[0013] The invention also relates to an isolated, soluble catalytically active polypeptide comprising the active conformation of the human Chk1 kinase or an active kinase analog thereof.

[0014] The invention encompasses both the polypeptide *per se* as well as salts thereof. As discussed in detail below, a high salt concentration (about 500 mM) in the buffer is used herein to prevent aggregation of peptide during purification and storage.

[0015] The invention also relates to a crystal structure of the human Chk1 kinase in the active conformation resolved to at least 2.5 (), preferably 2.0 (), more preferably 1.7 (). This structure provides a three-dimensional description of the target (human Chk1) for structure-based design of small molecule inhibitors thereof as therapeutic agents.

[0016] The invention further relates to an expression vector for producing catalytically active human Chk1 kinase in a host cell.

The invention further relates to a host cell stably transformed and transfected with a polynucleotide encoding of the human Chk1 kinase, or fragment thereof; or an active kinase analog thereof, in a manner allowing the expression of the human Chk1 kinase in the active configuration.

[0018] The present invention further discloses methods for screening candidate compounds using the molecular structure of the x-ray crystallography data to model the binding of candidate compounds.

[0019] The invention further provides a method for designing and screening potentially therapeutic compounds for the treatment of hyper-proliferative or diseases related to proliferation, including but not limited to cancer and HIV infection. The putative therapeutics can be screened for activities such as (1) potentiation of the cytotoxicity of DNA damaging agents such as synthetic or natural chemotherapeutic agents and ionizing or neutron radiation; (2) enhancement of the cytotoxicity of DNA synthesis inhibitors including antimetabolites, DNA chain terminators, or other mechanisms that would lead to the inhibition of DNA synthesis; (3) enhancement of the cytotoxicity of hypoxia as would occur within tumors due to a limited blood supply; and (4) inhibition of the ability of HIV to arrest cell cycle progression such as that induced by the VPR protein. Compounds that inhibit human Chk1 kinase activity or abrogate the G2 checkpoint can be used to treat or prevent the hyperproliferation associated with cancer and HIV.

[0020] The present invention provides methods for identifying potential inhibitors of the human Chk1 protein kinase by de novo design of novel drug candidate molecules that bind to and inhibit human Chk1 protein kinase activity, or that improve their potency. The x-ray crystallographic coordinates disclosed herein, allow generation of 3-dimensional models of the catalytic site and the drug binding site of the human Chk1 protein. De novo design comprises of the generation of molecules via the use of computer programs which build and link fragments or atoms into a site based upon steric and electrostatic complementarily, without reference to substrate analog structures. The drug design process begins after the structure of the target (human Chk1 kinase) is solved to at least a resolution of 2.5_. Refinement of the structure to a resolution of 2.0 Å or better with fixed water molecules in place provides more optimal conditions to undertake drug design.

[0021] The invention further provides a method for computational modeling of the kinase domain of human Chk1, such a model being useful in the design of compounds that interact with this domain. The method involves crystallizing the Chk1 kinase in the catalytically active configuration; resolving the x-ray structure of said active kinase, particularly the kinase domain and binding site of active Chk1; and applying the data generated from resolving the x-ray structure to a computer algorithm capable of generating a three dimensional model of the kinase domain and binding site suitable for use in designing molecules that will act as agonists or antagonists to the polypeptide. An iterative process can then be applied to various molecular structures using the computer-generated model to identify potential agonists or antagonists of the Chk1 kinase. Inhibitors of the kinase can serve as lead compounds for the design of potentially therapeutic compounds for the treatment of diseases or disorders associated with hyperproliferation or related to proliferation, such as cancer and HIV.

[0022] The invention further provides a process where the human Chk1 protein kinase is modified by deletion of the C-terminal portion of the protein so as to impart favorable physical characteristics of the resulting polypeptide. The kinase domain is suitable for analysis by nuclear magnetic resonance, high throughput screening, biochemical characterizations, x-ray crystallography, colorimetry and other diagnostic means. The most preferred deletion fragment extends from residue 1 to residue 289.

[0023] The invention further provides screening methods for use in the drug design process of potential agents to the human Chk1 protein kinase by *de novo* design of novel drug candidate molecules with potentially nanomolar potencies. The x-ray crystallographic coordinates disclosed based on the kinase domain of the human Chk1 protein will allow the generation of 3-dimensional models of the active binding sites of the human Chk1 protein.

The invention further provides a method for rapidly screening compounds to identify those compounds that inhibit Chk1 kinase or core structure for further Chk1 inhibitor design. The high throughput-screening assay is capable of being fully automated on robotic workstations. The assay may be radioactive. However, in a preferred embodiment the assay is a non-radioactive ELISA. In a more preferred embodiment, the assay is an ELISA that utilizes a novel antibody, rabbit anti-phosphosyntide, to specifically detect the product of the Chk1 kinase reaction in which biotin-syntide is the substrate. However, the basis of the assay includes the ability to use other substrates detectable by anti-phosphopeptide/ protein antibodies. The assay may be used to screen large collections of compound libraries to discover Chk1 kinase inhibitors and potential lead compounds for the development of Chk1 kinase selective anticancer compounds. The assay finds utility in the screening of other syntide substrate kinase reactions involving kinases of analogous activity to Chk1.

BRIEF DESCRIPTION OF THE DRAWINGS

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Figure 1. The G₂ DNA damage checkpoint mechanism in fission yeast (Furnari et al., *Science*, **277**: 1495-1497 (Sep. 5, 1997).

Figure 2. Sequence alignment of Chk1 kinase domains of human (hs) (SEQ ID NO: 2), mouse (mm) (SEQ ID NO: 18), Xenopus (xl) (SEQ ID NO: 19), fruit fly (dm) (SEQ ID NO: 20), C. elegans (ce) (SEQ ID NO: 21), S. cerevisiae (sc) (SEQ ID NO: 22), and S. pombe (sp) (SEQ ID NO: 23). Secondary structural elements of human Chk1 are shown above the alignment. The numbers of amino acids are shown on the right. Invariant residues among these species are in red and human Chk1 residues that also conserved in other species are in cyan.

Figure 3. The homology model of Chk1 kinase depicting the activation loop and its relationship to the catalytic loop and C helix. The Chk1 N and C-terminal lobes are shown. The fragments corresponding to the Chk1 C-helix are residues 50-58; the Chk1 catalytic loop are residues 129-132; and the Chk1 activation loop are residues 148-170.

Figure 4. The purification scheme for Chk1 kinase domain 1-289.

Figure 5. The structure of human Chk1 kinase domain identified using the crystal resolved to 1.7 Å. A ribbon diagram of the binary complex structure of Chk1 with AMP-PNP showing the secondary structural elements and the loops discussed in the text. The α -helices are shown in blue, the β -strands in cyan, the catalytic loop in orange, the activation loop in red. AMP-PNP and sulfate ion are shown as ball and stick models. The termini are denoted by N and C.

Figure 6. Catalytic site of Chk1. Cross section of the catalytic site of human Chk1 with AMP-PNP. Protein C α -ribbon representations are shown in purple for Chk1. The side chains of the catalytic site residues are shown as ball and stick models and are color-coded by atom type: carbon, green; nitrogen, blue; oxygen, red. The distances (_) along the dotted lines between the catalytic site residues are shown.

Figure 7. Molecular surface of the Chk1 with modeled CDC25C peptide. The molecular surface of Chk1 is colored as follows: basic side chains are shown in blue, acidic side chains in red, and non-polar side chains in violate. CDC25C peptide (residues 211-219) is shown as tick model and color-coded by atom type: carbon, green; nitrogen, blue; oxygen, red; sulfur, yellow.

Figure 8. Stereoview of representative electron density map. Figure 8A shows a stereoview of a representative portion of the experimental density at 1.5_ calculated to 3.0_ with the use of phases after solvent flattening. Superimposed on the density is the final refined model. Figure 8B shows a difference Fourier map calculated with native model-derived phases and coefficients IFO(AMP-PNP)I-IFO(native/apoenzyme)I to the diffraction of 1.7_ and contoured at 2.5_. The triphosphate moiety of AMP-PNP is disordered and is omitted from the model. No Mg²⁺ ions are observed.

Figure 9. Representation of the Chk1 binding sites, showing specifically the specificity pocket, the ATP binding site, and the Donor-Acceptor-Donor binding motif.

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Figure 10. The high throughput ELISA protocol.

Figure 11. The Chk1 crystal coordinates for the apoenzyme (isolated active Chk1 — Figure 11A) and the binary complex (Chk1 complexed with AMP-PNP, an ATP analog — Figure 11B) including the coordinates of the fixed water molecules.

DETAILED DESCRIPTION OF THE INVENTION

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[0026] DNA damage induces the arrest of the cell cycle at the G₂ checkpoint. The G₂ DNA damage checkpoint ensures maintenance of cell viability by delaying progression into mitosis in cells which have suffered genomic damage. The G₂ checkpoint is controlled by cell cycle checkpoint pathways which have been extensively studied (Hartwell, LH et al., Science, 246 (1989), supra; Nurse, P et al., Nat Med, 4 (10): 1103-6 (Oct 1998); Peng et al., Science, 277, (1997), supra; Furnari et al., Science, 277: 1495-1497 (Sep. 5, 1997); Zeng et al., Nature 395 (6701):507-510 (Oct. 1, 1998); Martinho et al., EMBO J, 17(24):7239-49 (Dec. 15, 1998); Nakajo et al., Dev. Biol. 207(2):432-44 (Mar. 15, 1999); Carr et al., Curr Biol., 5 (1995), supra). The model of the checkpoint mechanism in fission yeast is shown in Figure 1, Furnari, et al., Science, (1997), supra. As mentioned above, the regulation control system is highly conserved from yeast to humans.

[0027] DNA damage activates the checkpoint pathway by inhibiting the dephosphorylation of the mitotic kinase Cdc2 at the tyrosine-15 residue [Cdc2 (Y¹⁵-PO₄)], thereby inhibiting its mitotic initiating activity and arresting the cell cycle. This process is referred to as inhibitory phosphorylation. In order for mitosis to proceed, Cdc2 must be dephosphorylated, returning it to its active form. Phosphorylated Cdc2 is the substrate of Cdc25. Cdc25 is a dual specificity protein phosphatase that controls entry into mitosis by dephosphorylating the protein kinase Cdc2. In fission yeast, DNA damage also results in the activation of Rad3, a kinase related to the ATM/ATR proteins. Rad3 initiates the Chk1 response; the phosphorylation of Chk1 is a Rad3 dependent process (Martinho et al., *EMBO J*, 17 (1998), supra; Furnari et al., *Science*, 277 (1997), supra). Phosphorylated (active) Chk1 phosphorylates the mitotic inducer Cdc25 at the serine-216 residue of human Cdc25 [Cdc25 (S²¹⁶-PO₄)]. Phosphorylation of Cdc25 inhibits the function of the phosphatase in the dephosphorylated at the serine-216 residue and bound to members of the highly conserved and ubiquitously expressed family of 14-3-3 proteins. Prevention of serine-216 phosphorylation prevents 14-3-3 binding, perturbing mitotic timing and allowing cells to escape the G₂ checkpoint arrest induced by either unreplicated DNA or radiation induced damage.

[0028] A majority of currently accepted cancer treatments involve the induction of DNA damage including the

administration of anticancer agents, chemotherapeutic agents, and radiation therapy. Cancer cells frequently become resistant to such therapies. It is suspected that such resistance is related to the innate ability of the cancer cells to arrest and repair the damage induced. If the cancer cell was unable to arrest and repair, mitosis would proceed with the DNA damage intact. The downstream result would presumably be cell death as a result of the DNA damage.

[0029] Treatments that include a mechanism for abrogating the endogenous checkpoint pathway and repair process would presumably be more effective in killing cancer cells. As many cancer cells already lack a G_1 checkpoint control system, a therapy that involved the inhibition of the G_2 checkpoint would presumably force the cancer cells to proceed through mitosis without any feedback arrest and repair process. Hence, there is a clear utility for the inhibition of the activity of Chk1, a pivotal kinase in the G_2 checkpoint pathway. As many of the same events that regulate the G_2 arrest subsequent to DNA damage also regulate the S phase delay following DNA damage, the inhibition of Chk1 finds utility in the regulation of S phase as well.

[0030] The human Chk1 sequence of amino acids 1 to 476 is available through GenBank. Full length or segments of human Chk1 cDNA corresponding to codon 1-427, 1-265, and 1-289 were separately amplified by PCR. Each was tagged at its 3'-end with six histidine codons and cloned into an expression plasmid for protein production using a Baculovirus/insect cell expression system. The protein was expressed in insect Hi-5 cells and purified by a combination of ion-exchange and affinity column chromatography. It was found that a high concentration of salt (~500 mM levels) was required for keeping the purified Chk1 kinase domain from forming a precipitate.

[0031] The kinase activity of the hChk1 was determined by monitoring the ADP production through enzymatic actions of pyruvate kinase and lactate dehydrogenase. The Chk1 kinase domain containing amino acids 1-289 showed higher enzymatic activity than the full length protein. Unlike the other forms of Chk1 which have proven difficult to work with (isolate, purify, crystallize, etc), the 1-289 kinase domain form of the human Chk1 enzyme facilitated crystallography, enzyme characterization, and high throughput screening of inhibitors. In particular, the Chk1 kinase domain was used to determine its 3-dimensional structure, which provides unique structural information for inhibitor design for therapeutic development.

[0032] As used herein, the abbreviation 'hChk1' refers to the polynucleotide encoding the human effector check-point kinase serving as a DNA damage/replication checkpoint kinase. The nucleic acid sequence of the polynucleotide encoding the full length protein of human Chk1 was published in Science by Sanchez et al. (*Science*, 277 (5331): 1497-1501 (1997)) and published in GenBank on September 9, 1997 (AF016582). The nucleic acid sequence described therein is provided herein, shown in SEQ ID NO. 1. The corresponding peptide sequence of the full length protein is provided herein, shown in SEQ ID NO. 2. This peptide sequence was submitted to GenBank by Flaggs et al. on November 3, 1997 and released on December 13, 1997 (AF032874). The protein kinase was further described by Flaggs et al. in Current Biology (*Curr. Biol.*, 7(12):977-986, (1997)).

[0033] Using homology tools to examine the nucleotide and peptide sequence of Chk1, scientists have attempted to estimate the location of the kinase domain. However, the exact location of the catalytically active kinase domain has been difficult to experimentally determine, primarily as no one has ever reported isolating the kinase domain in its active configuration. Previous publications have indicated that the kinase domain extends from AA 16 to AA 264 (WO99/111795, published March 11, 1999, at page 7, line 3) of SEQ ID NO. 2.

[0034] We have found that the catalytic kinase domain begins between AA1 and 16 and terminates between AA265 and AA291-of-SEQ-ID-NO.-2.-We-further discovered that-vector-driven protein yield is dramatically increased when a fragment extending from AA1 to AA289 (dubbed KH289) is used.

[0035] There are 22 known amino acids but 64 possible permutations of nucleic acid triplets, called "codons". Many amino acids are specified by more than one codon, a phenomenon called degeneracy. Due to the degeneracy of the genetic code, there are many functionally equivalent nucleic acid sequences that can encode the same protein. The active human Chk1 kinase set forth in SEQ ID NO.2 can clearly be encoded by multiple nucleotide sequences and is not limited to the cDNA sequence set forth in SEQ ID NO. 1. For example, both UUU and UUC code for a phenylalanine while serine is encoded by UCU, UCC, UCA, UCG, AGU, and AGC [Molecular Biology of the Gene, 4th edition, Watson, J.D. et al., editors (1987) at pages 437-438]. Functionally equivalent sequences can readily be prepared using known methods such as modified primer PCR, site-directed mutagenesis, and chemical synthesis. Such functional equivalents are within the scope of this invention.

[0036] In the examples of the present invention, the full length form of human Chk1 protein kinase (AA 1-476) is referred to as KH476. Fragments thereof are identified by the amino acid sequence. For example, the human Chk1 kinase domain (AA 1-289) is referred to as KH289 Other kinase domain sequences are referred to by amino acid numbering in a similar manner.

A. Peptides, Proteins and Antibodies

[0037] As used herein, the terms "kinase" and "protein kinase" refer to enzymes that catalyze the transfer of a phosphate residue from a nucleoside triphosphate to an amino acid side chain in selected targets. The covalent phosphor-

ylation in turn regulates the activity of the target protein. In addition, phosphorylation frequently acts as the signal that triggers a particular process or reaction, playing an integral part in cellular regulation and control mechanisms. Clearly, inappropriate or unregulated phosphorylation can result in errors in cell signaling and the associated cell cycle and regulation processes. Most protein kinases are highly substrate specific.

[0038] As used herein, a peptide is said to be "isolated" or "purified" when it is substantially free of homologous cellular material or chemical precursors or other chemicals. The peptides of the present invention can be purified to homogeneity or other degrees of purity. The level of purification will be based on the intended use.

[0039] In some uses, "substantially free of cellular material" includes preparations of the peptide having less than about 30% (by dry weight) other proteins (i.e., contaminating protein), less than about 20% other proteins, less than about 10% other proteins, or less than about 5% other proteins. When the peptide is recombinantly produced, it can also be substantially free of culture medium, i.e., culture medium represents less than about 20% of the volume of the protein preparation.

[0040] The language "substantially free of chemical precursors or other chemicals" includes preparations of the peptide in which it is separated from chemical precursors or other chemicals that are involved in its synthesis. In one embodiment; the language "substantially free of chemical precursors or other chemicals" includes preparations of the kinase peptide having less than about 30% (by thy weight) chemical precursors or other chemicals, preferably less than about 20% chemical precursors or other chemicals, more preferably less than about 10% chemical precursors or other chemicals, or most preferably less than about 5% chemical precursors or other chemicals.

[0041] The isolated kinase described herein can be purified from cells that naturally express it, purified from cells that have been altered to express it (recombination), or synthesized using known protein synthesis methods. For example, a nucleic acid molecule encoding the protein kinase is cloned into an expression vector, the expression vector introduced into a host cell and the protein expressed in the host cell. The protein can then be isolated from the cells by an appropriate purification scheme using standard protein purification techniques. Many of these techniques are described in detail below.

[0042] The present invention also provides catalytically active variants of the peptides of the present invention, such as allelic/sequence variants of the peptides, non-naturally occurring recombinantly derived variants of the peptides, and orthologs and paralogs of the peptides. Such variants can be generated using techniques that are known by those skilled in the fields of recombinant nucleic acid technology and protein biochemistry.

[0043] Such variants can readily be identified/made using molecular techniques and the sequence information disclosed herein. Further, such variants can readily be distinguished from other peptides based on sequence and/or structural homology to the peptides of the present invention. The degree of homology/identity present will be based primarily on whether the peptide is a functional (active) variant or non-functional (inactive) variant, the amount of divergence present in the paralog family and the evolutionary distance between the orthologs.

[0044] To determine the percent identity of two amino acid sequences or two nucleic acid sequences, the sequences are aligned for optimal comparison purposes (e.g., gaps can be introduced in one or both of a first and a second amino acid or nucleic acid sequence for optimal alignment and non-homologous sequences can be disregarded for comparison purposes). In a preferred embodiment, the length of a reference sequence aligned for comparison purposes is at least 30%, 40%, 50%, 60%, 70%, 80%, or 90% or more of the length of the reference sequence. The amino acid residues or nucleotides at corresponding amino acid positions or nucleotide positions are then compared. When a position in the first sequence is occupied by the same amino acid residue or nucleotide as the corresponding position in the second sequence, then the molecules are identical at that position (as used herein amino acid or nucleic acid 'identity' is equivalent to amino acid or nucleic acid 'homology'). The percent identity between the two sequences is a function of the number of identical positions shared by the sequences, taking into account the number of gaps, and the length of each gap, which need to be introduced for optimal alignment of the two sequences.

[0045] The comparison of sequences and determination of percent identity and similarity between two sequences can be accomplished using a mathematical algorithm. (*Computational Molecular Biology*, Lesk, A.M., ed., Oxford University Press, New York, 1988; *Biocomputing: Informatics and Genome Projects*, Smith, D.W., ed., Academic Press, New York, 1993; *Computer Analysis of Sequence Data, Part 1*, Griffin, A.M., and Griffin, H.G., eds., Humana Press, New Jersey, 1994; *Sequence Analysis in Molecular Biology*, von Heinje, G., Academic Press, 1987; and *Sequence Analysis Primer*, Gribskov, M. and Devereux, J., eds., M Stockton Press, New York, 1991). In a preferred embodiment, the percent identity between two amino acid sequences is determined using the Needleman and Wunsch (*J. Mol. Biol.* (48):444-453 (1970)) algorithm which has been incorporated into commercially available computer programs, such as GAP in the GCG software package, using either a Blossom 62 matrix or a PAM250 matrix, and a gap weight of 16, 14, 12, 10, 8, 6, or 4 and a length weight of 1, 2, 3, 4, 5, or 6. In yet another preferred embodiment, the percent identity between two nucleotide sequences can be determined using the commercially available computer programs including the GAP program in the GCG software package (Devereux, J., et al., Nucleic Acids Res. 12(1):387 (1984)), the NWS gap DNA CMP matrix and a gap weight of 40, 50, 60, 70, or 80 and a length weight of 1, 2, 3, 4, 5, or 6. In another embodiment, the percent identity between two amino acid or nucleotide sequences is determined using the algorithm

of E. Meyers and W. Miller (CABIOS, 4:11-17 (1989)) which has been incorporated into commercially available computer programs, such as ALIGN (version 2:0), using a PAM120 weight residue table, a gap length penalty of 12 and a gap penalty of 4.

[0046] The nucleic acid and protein sequences of the present invention can further be used as a "query sequence" to perform a search against sequence databases to, for example, identify other family members or related sequences. Such searches can be performed using commercially available search engines, such as the NBLAST and XBLAST programs (version 2.0) of Altschul, et at. (*J. Mol. Biol.* 215:403-10 (1990)). Nucleotide searches can be performed with such programs to obtain nucleotide sequences homologous to the nucleic acid molecules of the invention. Protein searches can be performed with such programs to obtain amino acid sequences homologous to the proteins of the invention. To obtain gapped alignments for comparison purposes, Gapped BLAST can be utilized as described in Altschul et al. (*Nucleic Acids Res.* 25(17):3389-3402 (1997)).

[0047] Full-length clones comprising one of the peptides of the present invention can readily be identified as having complete sequence identity to one of the kinases of the present invention as well as being encoded by the same genetic locus as the kinase provided herein.

[0048] Allelic variants of a peptide can readily be identified as having a high degree (significant) of sequence homology/identity to at least a portion of the peptide as well as being encoded by the same genetic locus as the kinase peptide provided herein. As used herein, two proteins (or a region of the proteins) have significant homology when the amino acid sequences are typically at least about 70-75%, 80-85%, and more typically at least about 90-95% or more homologous. A significantly homologous amino acid sequence, according to the present invention, will be encoded by a nucleic acid sequence that will hybridize to a peptide encoding nucleic acid molecule under siringent conditions as more fully described below.

[0049] Paralogs of a peptide can readily be identified as having some degree of significant sequence homology/identity to at least a portion of the kinase peptide, as being encoded by a gene from Drosophila, and as having similar activity or function. Two proteins will typically be considered paralogs when the amino acid sequences are typically at least about 70-75%, 80-85%, and more typically at least about 90-95% or more homologous through a given region or domain. Such paralogs will be encoded by a nucleic acid sequence that will hybridize to a kinase peptide encoding nucleic acid molecule under stringent conditions as more fully described below.

[0050] Orthologs of a kinase peptide can readily be identified as having some degree of significant sequence homology/identity to at least a portion of the kinase peptide as well as being encoded by a gene from another organism. Preferred orthologs will be isolated from mammals, preferably human, for the development of human therapeutic targets and agents, or other invertebrates, particularly insects of economical/agriculture importance, e.g. members of the Lepidopteran and Coleopteran orders, for the development of insecticides and insecticidal targets. Such orthologs will be encoded by a nucleic acid sequence that will hybridize to a kinase peptide encoding nucleic acid molecule under moderate to stringent conditions, as more fully described below, depending on the degree of relatedness of the two organisms yielding the proteins.

[0051] Non-naturally occurring variants of the kinases of the present invention can readily be generated using recombinant techniques. Such variants include, but are not limited to deletions, additions and substitutions in the amino acid sequence of the kinase. For example, one class of substitutions are conserved amino acid substitution. Such substitutions are those that substitute a given amino acid in a kinase peptide by another amino acid of like characteristics. Typically seen as conservative substitutions are the replacements, one for another, among the aliphatic amino acids Ala, Val, Leu, and Ile; interchange of the hydroxyl residues Ser and Thr; exchange of the acidic residues Asp and Glu; substitution between the amide residues Asn and Gln; exchange of the basic residues Lys and Arg; and replacements among the aromatic residues Phe, Tyr. Guidance concerning which amino acid changes are likely to be phenotypically silent are found in Bowie et al., Science 247:1306-1310 (1990).

[0052] Variant kinases can be fully functional or can lack function in one or more activities. Fully functional variants typically contain only conservative variation or variation in non-critical residues or in non-critical regions. Functional variants can also contain substitution of similar amino acids, which result in no change or an insignificant change in function. Alternatively, such substitutions may positively or negatively affect function to some degree.

[0053] Non-functional variants typically contain one or more non-conservative amino acid substitutions, deletions, insertions, inversions, or truncation or a substitution, insertion, inversion, or deletion in a critical residue or critical region.

[0054] Amino acids that are essential for function can be identified by methods known in the art, such as site-directed mutagenesis or alanine-scanning mutagenesis (Cunningham *el al.*, *Science 244*:1081-1085 (1989)). The latter procedure introduces single alanine mutations at every residue in the molecule. The resulting mutant molecules are then tested for biological activity such as receptor binding or *in vitro* proliferative activity. Sites that are critical for binding can also be determined by structural analysis such as x-ray crystallography, nuclear magnetic resonance or photoaffinity labeling (Smith *et al.*, *J. Mol. Biol. 224*:899-904 (1992); de Vos *et al. Science 255*:306-312 (1992)). Accordingly, the protein kinases of the present invention also encompass derivatives or analogs in which a substituted amino acid resi-

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due is not one encoded by the genetic code; in which a substituent group is included; in which the mature polypeptide is fused with another compound, such as a compound to increase the half-life of the polypeptide (for example, polyeth-ylene glycol); or in which the additional amino acids are fused to the mature polypeptide, such as a leader or secretory sequence or a sequence for purification of the mature polypeptide or a pro-protein sequence.

[0055] The present invention further provides for functional, active fragments of the Chk1 kinase domain. As used herein, a fragment comprises at least 8 or more contiguous amino acid residues from the protein kinase. Such fragments can be chosen based on the ability to retain one or more of the biological activities of the kinase or could be chosen for the ability to perform a function, e.g. act as an immunogen. Particularly important fragments are catalytically activate fragments, peptides which are, for example about 8 or more amino acids in length. Such fragments will typically comprise a domain or motif of the kinase, e.g., active site or binding site. Further fragments contemplated by the present invention include, but are not limited to, domain or motif containing fragments, soluble peptide fragments, and fragments containing immunogenic structures. Predicted domains and functional sites available to those of skill in the art (e.g., by PROSITE analysis).

[0056] Polypeptides often contain amino acids other than the 20 amino acids commonly referred to as the 20 naturally-occurring amino acids. Further, many amino acids, including the terminal amino acids, may be modified by natural processes, such as processing and other post-translational modifications, or by chemical modification techniques known in the art. Common modifications that occur naturally in polypeptides are described in basic texts, detailed monographs, and the research literature, and they are known to those of skill in the art.

[0057] Known modifications include, but are not limited to, acetylation, acylation, ADP-ribosylation, amidation, covalent attachment of flavin, covalent attachment of a heme moiety, covalent attachment of a nucleotide derivative, covalent attachment of a lipid or lipid derivative, covalent attachment of phosphotidylinositol, cross-linking, cyclization, disulfide bond formation, demethylation, formation of covalent crosslinks, formation of cystine, formation of pyroglutamate, formylation, gamma carboxylation, glycosylation, GPI anchor formation, hydroxylation, iodination, methylation, myristoylation, oxidation, proteolytic processing, phosphorylation, phenylation, racemization, selenoylation, sulfation, transfer-RNA mediated addition of amino acids to proteins such as arginylation, and ubiquitination. Such modifications are known to those of skill in the art and have been described in great detail in the scientific literature. Several particularly common modifications, glycosylation, lipid attachment; sulfation, gamma-carboxylation of glutamic acid residues, hydroxylation and ADP-ribosylation, for instance, are described in most basic texts, such as *Proteins - Structure and Molecular Properties*, 2nd Ed., T.E. Creighton, W. H. Freeman and Company, New York (1993). Many detailed reviews are available on this subject; such as by Wold, F., *Posttranslational Covalent Modification of Proteins*, B.C. Johnson, Ed., Academic Press, New York 1-12 (1983); Seifter *et al.* (*Meth. Enzymol. 182*: 626-646 (1990)) and Rattan *et al.* (*Ann. N.Y. Acad Sci. 663*:48-62 (1992)).

[0058] The peptides of the present invention can be attached to heterologous sequences to form chimeric or fusion proteins. Such chimeric and fission proteins comprise a peptide operatively linked to a heterologous protein having an amino acid sequence not substantially homologous to the kinase peptide. "Operatively linked" indicates that the peptide and the heterologous protein are fused in-frame. The heterologous protein can be fused to the N-terminus or C-terminus of the kinase peptide. The two peptides linked in a fusion peptide are typically derived from two independent sources, and therefore a fusion peptide comprises two linked peptides not normally found linked in nature. The two peptides may be from the same or different genome.

[0059] In some uses, the fusion protein does not affect the activity of the peptide *per se*. For example, the fusion protein can include, but is not limited to, enzymatic fusion proteins, for example beta-galactosidase fusions, yeast two-hybrid GAL fusions, poly-His fusions, MYC-tagged, HI-tagged and Ig fusions. Such fusion proteins, particularly poly-His fusions, can facilitate the purification of recombinant kinase peptide. In certain host cells (e.g., mammalian host cells), expression and/or secretion of a protein can be increased by using a heterologous signal sequence.

[0060] A chimeric or fusion protein can be produced by standard recombinant DNA techniques. For example, DNA fragments coding for the different protein sequences are ligated together in-frame in accordance with conventional techniques. In another embodiment; the fusion gene can be synthesized by conventional techniques including automated DNA synthesizers. Alternatively, PCR amplification of gene fragments can be carried out using anchor primers which give rise to complementary overhangs between two consecutive gene fragments which can subsequently be annealed and re-amplified to generate a chimeric gene sequence (see Ausubel et al., Current Protocols in Molecular Biology, 1992). Moreover, many expression vectors are commercially available that already encode a fusion moiety (e.g., a GST protein). A kinase peptide-encoding nucleic acid can be cloned into such an expression vector such that the fusion moiety is linked in-frame to the kinase peptide.

[0061] Herein, the term 'antibody' refers to a polypeptide or group of polypeptides which are comprised of at least one antibody combining site or binding domain, said binding domain or combining site formed from the folding of variable domains of an antibody molecule to form three dimensional binding spaces with an internal surface shape and charge distribution complementary to the features of an antigen epitope. The term encompasses immunoglobulin molecules and immunologically active portions of immunoglobulin molecules, such as molecules that contain an antibody

combining site or paratope. Exemplary antibody molecules are intact immunoglobulin molecules, substantially intact immunoglobulin molecules and portions of an immunoglobulin molecule, including those known in the art as Fab, FabB, F(abB)₂ and F(v).

B. Nucleic Acids and Polynucleotides

[0062] The present invention provides isolated nucleic acid molecules that encode the functional, active kinases of the present invention. Such nucleic acid molecules will consist of, consist essentially of, or comprise a nucleotide sequence that encodes one of the kinase peptides of the present invention, an allelic variant thereof, or an ortholog or paralog thereof.

[0063] As used herein, an "isolated" nucleic acid molecule is one that is separated from other nucleic acid present in the natural source of the nucleic acid. Preferably, an "isolated" nucleic acid is free of sequences which naturally flank the nucleic acid (i.e., sequences located at the 5' and 3' ends of the nucleic acid) in the genomic DNA or cDNA of the organism from which the nucleic acid is derived. However, there can be some flanking nucleotide sequences, for example up to about 5KB, particularly contiguous peptide encoding sequences and peptide encoding sequences within the same gene but separated by introns in the genomic sequence. The important point is that the nucleic acid is isolated from remote and unimportant flanking sequences such that it can be subjected to the specific manipulations described herein such as recombinant expression, preparation of probes and primers, and other uses specific to the nucleic acid sequences.

[0064] Moreover, an "isolated" nucleic acid molecule, such as a cDNA molecule, can be substantially free of other cellular material, or culture medium when produced by recombinant techniques, or chemical precursors or other chemicals when chemically synthesized. However, the nucleic acid molecule can be fused to other coding or regulatory sequences and still be considered isolated.

[0065] For example, recombinant DNA molecules contained in a vector are considered isolated. Further examples of isolated DNA molecules include recombinant DNA molecules maintained in heterologous host cells or purified (partially or substantially) DNA molecules in solution. Isolated RNA molecules include *in vivo* or *in vitro* RNA transcripts of the isolated DNA molecules of the present invention. Isolated nucleic acid molecules according to the present invention further include such molecules produced synthetically.

[0066] The preferred classes of nucleic acid molecules that are comprised of the nucleotide sequences of the present are the full-length cDNA molecules and genes and genomic clones since some of the nucleic acid molecules provided in SEQ ID NO. 1 are fragments of the complete gene that exists in nature. A brief description of how various types of these nucleic acid molecules can be readily made/isolated is provided herein.

[0067] Full-length genes may be cloned from known sequence using any one of a number of methods known in the art. For example, a method which employs XL-PCR (Perkin-Elmer, Foster City, Calif.) to amplify long pieces of DNA may be used. Other methods for obtaining full-length sequences are known in the art.

[0068] The isolated nucleic acid molecules can encode the functional, active kinase plus additional amino or carboxyl-terminal amino acids, such as those that facilitate protein trafficking, prolong or shorten protein half-life or facilitate manipulation of a protein for assay or production, among other things. The isolated nucleic acid molecules include, but are not limited to, the sequence encoding the active kinase alone or in combination with coding sequences, such as a leader or secretory sequence (eg., a pre-pro or pro-protein sequence), the sequence encoding the active kinase, with or without the additional coding sequences, plus additional non-coding sequences, for example introns and non-coding 5' and 3' sequences such as transcribed but non-translated sequences that play a role in transcription, mRNA processing (including splicing and polyadenylation signals), ribosome binding and stability of mRNA. In addition, the nucleic acid molecule may be fused to a marker sequence encoding, for example, a peptide that facilitates purification.

[0069] Isolated nucleic acid molecules can be m the form of RNA, such as mRNA, or m the form DNA, including cDNA and genomic DNA, obtained by cloning or produced by chemical synthetic techniques or by a combination thereof The nucleic acid, especially DNA, can be double-stranded or single-stranded. Single-stranded nucleic acid can be the coding strand (sense strand) or the non-coding strand (anti-sense strand).

[0070] The invention further provides nucleic acid molecules that encode functional fragments or variants of the active kinases of the present invention. Such nucleic acid molecules may be naturally occurring, such as allelic variants (same locus), paralogs (different locus), and orthologs (different organism), or may be constructed by recombinant DNA methods or by chemical synthesis. Such non-naturally occurring variants may be made by mutagenesis techniques, including those applied to nucleic acid molecules, cells, or organisms. Accordingly, as discussed above, the variants can contain nucleotide substitutions, deletions, inversions and insertions. Variation can occur in either or both the coding and non-coding regions. The variations can produce both conservative and non-conservative amino acid substitutions.

[0071] A fragment comprises a contiguous nucleotide sequence greater than 12 or more nucleotides. Further, a fragment could be at least 30, 40, 50, 100, 250 or 500 nucleotides in length. The length of the fragment will be based

on its intended use. For example, the fragment can encode epitope bearing regions of the peptide, or can be useful as DNA probes and primers. Such fragments can be isolated using the known nucleotide sequence to synthesize an oligonucleotide probe. A labeled probe can then be used to screen a cDNA library, genomic DNA library, or mRNA to isolate nucleic acid corresponding to the coding region. Further, primers can be used in PCR reactions to clone specific regions of gene.

[0072] A probe/primer typically comprises substantially a purified oligonucleotide or oligonucleolide pair. The oligonucleotide typically comprises a region of nucleotide sequence that hybridizes under stringent conditions to at least about 12, 20, 25, 40, 50 or more consecutive nucleotides.

[0073] Orthologs, homologs, and allelic variants can be identified using methods known in the art. As described above, these variants comprise a nucleotide sequence encoding a peptide that is typically 60-65%, 70-75%, 80-85%, and more typically at least about 90-95% or more homologous to the nucleotide sequence provided in SEQ ID NO. 1 or a fragment of this sequence. Such nucleic acid molecules can readily be identified as being able to hybridize under moderate to stringent conditions, to the nucleotide sequence shown in SEQ ID NO. 1 or a fragment of the sequence.

[0074] As used herein, the term "hybridizes under stringent conditions" is intended to describe conditions for hybridization and washing under which nucleotide sequences encoding a peptide at least 50-55% homologous to each other typically remain hybridized to each other. The conditions can be such that sequences at least about 65%, at least about 70%, or at least about 75% or more homologous to each other typically remains hybridized to each other. Such stringent conditions are known to those skilled in the art and can be found in *Current Protocols in Molecular Biology*, John Wiley & Sons, N.Y. (1989), 6.3.1-6.3.6. One example of stringent hybridization conditions are hybridization in 6X sodium chloride/sodium citrate (SSC) at about 45C, followed by one or more washes in 0.2 X SSC, 0.1% SDS at 50-65C.

[0075] The nucleic acid molecules of the present invention are useful for probes, primers, chemical intermediates, and in biological assays. The nucleic acid molecules are useful as a hybridization probe for cDNA and genomic DNA to isolate full-length cDNA and genomic clones encoding the peptide described herein and to isolate cDNA and genomic clones that correspond to variants (alleles, orthologs, etc.) producing the same or related peptides described herein.

[0076] The nucleic acid molecules are also useful as primers for PCR to amplify any given region of a nucleic acid molecule and are useful to synthesize antisense molecules of desired length and sequence.

[0077] The nucleic acid molecules are also useful for constructing recombinant vectors. Such vectors include expression vectors that express a portion of; or all of, the peptide sequences. Vectors also include insertion vectors, used to integrate into another nucleic acid molecule sequence, such as into the cellular genome, to alter *in situ* expression of a gene and/or gene product. For example, an endogenous coding sequence can be replaced via homologous recombination with all or part of the coding region containing one or more specifically introduced mutations.

[0078] The nucleic acid molecules are also useful for expressing antigenic portions of the proteins.

[0079] The nucleic acid molecules are also useful as probes for determining the chromosomal positions of the nucleic acid molecules by means of *in situ* hybridization methods.

[0080] The nucleic acid molecules are also useful for designing ribozymes corresponding to all, or a part, of the mRNA produced from the nucleic acid molecules described herein.

[0081] The nucleic acid molecules are also useful for constructing host cells expressing a part, or all, of the nucleic acid molecules and peptides.

[0082] The nucleic acid molecules are also useful for constructing transgenic animals expressing all, or a part, of the nucleic acid molecules and peptides.

[0083] The nucleic acid molecules are also useful for making vectors that express part, or all, of the peptides.

[0084] The nucleic acid molecules are also useful as hybridization probes for determining the presence, level, form and distribution of nucleic acid expression. Accordingly, the probes can be used to detect the presence of; or to determine levels of, a specific nucleic acid molecule in cells, tissues, and in organisms. The nucleic acid whose level is determined can be DNA or RNA. Accordingly, probes corresponding to the peptides described herein can be used to assess expression and/or gene copy number in a given cell, tissue, or organism. These uses are relevant for diagnosis of disorders involving an increase or decrease in kinase protein expression relative to normal results.

[0085] In vitro techniques for detection of mRNA include Northern hybridizations and in situ hybridizations. In vitro techniques for detecting DNA includes Southern hybridizations and in situ hybridization.

[0086] Probes can be used as a part of a diagnostic test kit for identifying cells or tissues that express a kinase protein, such as by measuring a level of a receptor-encoding nucleic acid in a sample of cells from a subject e.g., mRNA or genomic DNA, or determining if a receptor gene has been mutated.

C. Vectors and Host Cells

[0087] The invention also provides vectors containing the nucleic acid molecules described herein. The term "vector" refers to a vehicle, preferably a nucleic acid molecule, that can transport the nucleic acid molecules. When the vector

tor is a nucleic acid molecule, the nucleic acid molecules are covalently linked to the vector nucleic acid. With this aspect of the invention, the vector includes a plasmid, single or double stranded phage, a single or double stranded RNA or DNA viral vector, or artificial chromosome, such as a BAC, PAC, YAC, OR MAC. Various expression vectors can be used to express polynucleotide encoding the active hChk1 kinase.

[0088] A vector can be maintained in the host cell as an extrachromosomal element where it replicates and produces additional copies of the nucleic acid molecules. Alternatively, the vector may integrate into the host cell genome and produce additional copies of the nucleic acid molecules when the host cell replicates.

[0089] The invention provides vectors for the maintenance (cloning vectors) or vectors for expression (expression vectors) of the nucleic acid molecules. The vectors can function in prokaryotic or eukaryotic cells or in both (shuttle vectors).

[0090] Expression vectors contain cis-acting regulatory regions that are operably linked in the vector to the nucleic acid molecules such that transcription of the nucleic acid molecules is allowed in a host cell. The nucleic acid molecules can be introduced into the host cell with a separate nucleic acid molecule capable of affecting transcription. Thus, the second nucleic acid molecule may provide a trans-acting factor interacting with the cis-regulatory control region to allow transcription of the nucleic acid molecules from the vector. Alternatively, a trans-acting factor may be supplied by the host cell. Finally, a trans-acting factor can be produced from the vector itself. It is understood, however, that in some embodiments, transcription and/or translation of the nucleic acid molecules can occur in a cell-free system.

[0091] The regulatory sequence to which the nucleic acid molecules described herein can be operably linked include promoters for directing mRNA transcription. These include, but are not limited to, the left promoter from bacteriophage λ , the lac, TRP, and TAC promoters from *E. coli*, the early and late promoters from SV40, the CMV immediate early promoter, the adenovirus early and late promoters, and retrovirus long-terminal repeats.

[0092] In addition to control regions that promote transcription, expression vectors may also include regions that modulate transcription, such as repressor binding sites and enhancers. Examples include the SV40 enhancer, the cytomegalovirus immediate early enhancer, polyoma enhancer, adenovirus enhancers, and retrovirus LTR enhancers. [0093] In addition to containing sites for transcription initiation and control, expression vectors can also contain sequences necessary for transcription termination and, in the transcribed region a ribosome binding site for translation. Other regulatory control elements for expression include initiation and termination codons as well as polyadenylation signals. The person of ordinary skill in the art would be aware of the numerous regulatory sequences that are useful in expression vectors. Such regulatory sequences are described, for example, in Sambrook et al., (Molecular Cloning: A Laboratory Manual. 2nd ed., Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY, (1989)).

[0094] A variety of expression vectors can be used to express a nucleic acid molecule. Such vectors include chromosomal, episomal, and virus-derived vectors, for example vectors derived from bacterial plasmids, from bacteriophage, from yeast episomes, from yeast chromosomal elements, including yeast artificial chromosomes, from viruses such as baculoviruses, papovaviruses such as SV40, Vaccinia viruses, adenoviruses, poxviruses, pseudorabies viruses, and retroviruses. Vectors may also be derived from combinations of these sources such as those derived from plasmid and bacteriophage genetic elements, eg. cosmids and phagemids. Appropriate cloning and expression vectors for prokaryotic and eukaryotic hosts are described in Sambrook et al., Molecular Cloning: A Laboratory Manual. 2nd ed., Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY, (1989).

[0095] The regulatory sequence may provide constitutive expression in one or more host cells (i.e. tissue specific) or may provide for inducible expression in one or more cell types such as by temperature, nutrient additive, or exogenous factor such as a hormone or other ligand. A variety of vectors providing for constitutive and inducible expression in prokaryotic and eukaryotic hosts are known to those of ordinary skill in the art.

[0096] The nucleic acid molecules can be inserted into the vector nucleic acid by well-known methodology. Generally, the DNA sequence that will ultimately be expressed is joined to an expression vector by cleaving the DNA sequence and the expression vector with one or more restriction enzymes and then ligating the fragments together. Procedures for restriction enzyme digestion and ligation are known to those of ordinary skill in the art.

[0097] The vector containing the appropriate nucleic acid molecule can be introduced into an appropriate host cell for propagation or expression using well-known techniques. Bacterial cells include, but are not limited to, *E. coli, Streptomyces, and Salmonella typhimurium*. Eukaryotic cells include, but are not limited to, yeast, insect cells such as *Drosophila*, animal cells such as COS and CHO cells, and plant cells.

As described herein, it may be desirable to express a peptide of the present invention as a fusion protein. Accordingly, the invention provides fusion vectors that allow for the production of such peptides. Fusion vectors can increase the expression of a recombinant protein, increase the solubility of the recombinant protein, and aid in the purification of the protein by acting for example as a ligand for affinity purification. A proteolytic cleavage site may be introduced at the junction of the fusion moiety so that the desired peptide can ultimately be separated from the fusion moiety. Proteolytic enzymes include, but are not limited to, factor Xa, thrombin, and enterokinase. Typical fusion expression vectors include pGEX (Smith *et al.*, *Gene 67*:31-40 (1988)), pMAL (New England Biolabs, Beverly, MA) and pRIT5 (Pharmacia, Piscataway, NJ) which fuse glutathione S-transferase (GST), maltose E binding protein, or protein A.

respectively, to the target recombinant protein. Examples of suitable inducible non-fusion *E. coli* expression vectors include pTrc (Amann et al., Gene 69:301-315 (1988)) and pET 11d (Studier et al., Gene Expression Technology: Methods in Enzymology 185:60-89 (1990)).

[0099] Recombinant protein expression can be maximized in a host bacteria by providing a genetic background wherein the host cell has an impaired capacity to proteolytically cleave the recombinant protein. (Gottesman, *S., Gene Expression Technology: Methods in Enzymology* 185, Academic Press, San Diego, California (1990) 119-128). Alternatively, the sequence of the nucleic acid molecule of interest can be altered to provide preferential codon usage for a specific host cell, for example *E. coli*. (Wada *et al.*, *Nucleic Acids Res. 20*:2111-2118 (1992)).

[0100] The nucleic acid molecules can also be expressed by expression vectors that are operative in yeast Examples of vectors for expression in yeast e.g., *S. cerevisiae* include pYepSec1 (Baldari, *et al.*, *EMBO J. 6*:229-234 (1987)), pMFa (Kurjan *et al.*, *Cell 30*:933-943(1982)), pJRY88 (Schultz *et al.*, *Gene 54*:113-123 (1987)), and pYES2 (Invitrogen Corporation, San Diego, CA).

[0101] The nucleic acid molecules can also be expressed in insect cells using, for example, baculovirus expression vectors. Baculovirus vectors available for expression of proteins in cultured insect cells (e.g., Sf 9 cells) include the pAc series (Smith *et al.*, *Mol. Cell Biol. 3*:2156-2165 (1983)) and the pVL series (Lucklow *et al.*, *Virology 170*:31-39 (1989)).

[0102] In certain embodiments of the invention, the nucleic acid molecules described herein are expressed in mammalian cells using mammalian expression vectors. Examples of mammalian expression vectors include pCDM8 (Seed, B. *Nature 329*:840(1987)) and pMT2PC (Kanfman *et al.*, *EMBO J. 6*:187-195 (1987)).

The expression vectors listed herein are provided by way of example only of the well-known vectors available to those of ordinary skill in the art that would be useful to express the nucleic acid molecules. Preferred vectors include the pET28a (Novagen, Madison, WI), pAcSG2 (Pharmingen, San Diego, CA), and pFastBac (Life Technologies, Gaithersburg. MD). The person of ordinary skill in the art would be aware of other vectors suitable for maintenance prepagation or expression of the nucleic acid molecules described herein. These are found for example in Sambrook, J., Fritsh, E. F., and Maniatis, T. *Molecular Cloning: A Laboratory Manual. 2nd, ed, Cold Spring Harbor Laboratory*, Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY, 1989.

[0104] The invention also encompasses vectors in which the nucleic acid sequences described herein are cloned into the vector in reverse orientation, but operably linked to a regulatory sequence that permits transcription of antisense RNA. Thus, an antisense transcript can be produced to all, or to a portion, of the nucleic acid molecule sequences described herein, including both coding and non-coding regions. Expression of this antisense RNA is subject to each of the parameters described above in relation to expression of the sense RNA (regulatory sequences stitutive or inducible expression, tissue-specific expression).

[0105] The invention also relates to recombinant host cells containing the vectors described herein. Host cells therefore include prokaryotic cells, lower eukaryotic cells such as yeast, other eukaryotic cells such as insect cells; and higher eukaryotic cells such as mammalian cells. Preferred host cells of the instant invention include *E. coli* and Sf9.

[0106] The recombinant host cells are prepared by introducing the vector constructs described herein into the cells by techniques readily available to the person of ordinary skill in the art. These include, but are not limited to, calcium phosphate transfection, DEAE-dextran-mediated transfection, cationic lipid-mediated transfection, electroporation, transduction, infection, lipofection, and other techniques such as those found in Sambrook, et al. (Molecular Cloning: A Laboratory Manual. 2nd, ed, Cold Spring Harbor Laboratory, Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY, 1989).

[0107] Host cells can contain more than one vector. Thus, different nucleotide sequences can be introduced on different vectors of the same cell. Similarly, the nucleic acid molecules can be introduced either alone or with other nucleic acid molecules that are not related to the nucleic acid molecules such as those providing trans-acting factors for expression vectors. When more than one vector is introduced into a cell, the vectors can be introduced independently, co-introduced or joined to the nucleic acid molecule vector.

[0108] In the case of bacteriophage and viral vectors, these can be introduced into cells as packaged or encapsulated virus by standard procedures for infection and transduction. Viral vectors can be replication-competent or replication-defective. In the case in which viral replication is defective, replication will occur in host cells providing functions that complement the defects.

[0109] Vectors generally include selectable markers that enable the selection of the subpopulation of cells that contain the recombinant vector constructs. The marker can be contained in the same vector that contains the nucleic acid molecules described herein or may be on a separate vector. Markers include tetracycline or ampicillin-resistance genes for prokaryotic host cells and dihydrofolate reductase or neomycin resistance for eukaryotic host cells. However, any marker that provides selection for a phenotypic trait will be effective.

While the active protein kinases can be produced in bacteria, yeast, mammalian cells, and other cells under the control of the appropriate regulatory sequences, cell- free transcription and translation systems can also be used to produce these proteins using RNA derived from the DNA constructs described herein.

[0111] Where secretion of the peptide is desired, appropriate secretion signals are incorporated into the vector. The

signal sequence can be endogenous to the peptides or heterologous to these peptides.

[0112] It is also understood that depending upon the host cell in recombinant production of the peptides described herein, the peptides can have various glycosylation patterns, depending upon the cell, or maybe non-glycosylated as when produced in bacteria in addition, the peptides may include an initial modified methionine in some cases as a result of a host-mediated process.

[0113] The recombinant host cells expressing the peptides described herein have a variety of uses. First, the cells are useful for producing a kinase protein or peptide that can be further purified to produce desired amounts of kinase protein or fragments. Thus, host cells containing expression vectors are useful for peptide production.

[0114] Host cells are also useful for conducting cell-based assays involving the kinase protein or kinase protein fragments. Thus, a recombinant host cell expressing a native kinase protein is useful for assaying compounds that stimulate or inhibit kinase protein function.

[0115] Host cells are also useful for identifying kinase protein mutants in which these functions are affected. If the mutants naturally occur and give rise to a pathology, host cells containing the mutations are useful to assay compounds that have a desired effect on the mutant kinase protein (for example, stimulating or inhibiting function) which may not be indicated by their effect on the native kinase protein.

[0116] The following examples are provided for illustration purposes.

Examples

1. Identification of the Catalytic Domain Sequence

[0117] From the complete protein sequence for the human checkpoint effector kinase (Chk1, 476 residues) available through GenBank, using sequence alignment and structures for other kinases, a homology model was devised for the kinase domain of the Chk1 protein (See Figure 3).

[0118] All protein kinases utilize ATP to phosphorylate their substrates, involving the transfer of a gamma phosphate to a substrate hydroxyl group. Each kinase binds ATP with its own strength, a property that is correlated by measuring K₁/IC50. The ATP molecule consists of adenine, ribose and triphosphate moleties. Each of these moleties bind to the protein in the ATP binding site (or ATP pokket). The adenine moiety always binds to the protein backbone by formation of two or three hydrogen bonds. The ribose moiety forms one to two hydrogen bonds with the protein side chains of amino acids that lay outside of the ATP pocket. The tri-phosphate moiety interacts with those catalytic amino acids of the kinase that are generally consistent across the whole protein kinase family. There is a limited specificity for each kinase within ATP binding groove. This region is referred to as the specificity pocket. Using the homology model, a schematic of the Chk1 binding site was developed, identifying the ATP binding site, the donor-acceptor-donor binding motif and the specificity pocket (See Figure 9). This binding site is the target for inhibitor development, e.g. the development of compounds or molecules that bind to this site to the extent that the kinase activity of the Chk1 protein is blocked or inhibited. The black and red color in Figure 9 represents the ATP binding groove; note, Ser 147 can contribute to the binding of inhibitor. The area designated by the blue color represents the region outside of the ATP pocket that can be used for enhancement of the specificity of binding. Finally, the area in pink represents the 'specificity pocket', that region that is very different from one protein to another. This site does not contribute to the ATP binding but can be used for the design of specific inhibitors. In other words, by utilizing that region of the Chk 1 binding site that is unique to Chk1 (the specificity pocket), one may design compounds that specifically inhibit Chk1 without also inhibiting the various other kinase molecules that may not be targets of the inhibition therapy.

[0119] Analysis of the C-termini of the kinase suggested that amino acids beyond residue 265 would enhance high level expression and/or maintain the appropriate crystal structure. The homology model showed this region to be flexible, such that ending the kinase domain construct within this region can prevent the disruption of potential secondary structures. Specifically, cleaving the Chk1 protein anywhere between amino acid residues 263 and 265 would result in the destruction of helical interactions at the distal end. The homology model further predicted that the kinase segment should extend to at least residue 272 to 275 and may be further extended to residue 289-291.

[0120] In addition, including the extended region in the construct prevents the C-terminal histidine tag from interacting with the kinase domain, making it accessible for affinity chromatography. Based on these analyses, construct KH289 was designed for the expression of Chk1 kinase domain of residue 1-289 with 6xHis-tag at its C-terminus. A corresponding construct without the 6xHis-tag was also made. Two other constructs were designed based on the homology model: (1) kinase domain of residues 1-210 (KH210) and (2) kinase domain of residues 1-248 (KH248).

55 2. Cloning

[0121] Human Chk1 cDNA was cloned by PCR using Vent polymerase (New England Biolabs, Beverly, MA) from human thymus and testis Marathon-Ready cDNA (Clontech, Palo Alto, CA) with primers synthesized (Genset, LaJolla,

A) based on the published sequence [SEQ ID NO. 1] (GenBank Accession number AF1016582) [Sanchez, Science (1997), supra.], following the instruction from the venders. Two overlapping sequences were amplified independently, one contained the sequence of nucleotides 35-830 of SEQ ID NO.1, and the other contained the sequence of nucleotides 678-1480 of SEQ ID NO.1. These overlapping sequences cover the whole coding sequence of Chk1 plus 16 base-pairs (bps) of 3'- untranslational region. The cDNA of 35-830 encodes the kinase domain of residues 1-265.

[0122] The PCR oligonucleotide primer sequences are listed in Table 1. Restriction sites for cloning, codons for 6xHis-tag, and the stop codon were engineered in the PCR primers. Restriction site Stul preceded Ncol site which overlapped the initiation codon. Sacl site followed the stop codon. When included, codons for 6xHis-tag preceded the stop codon, so that an expressed protein would have a 6xHis-tag at its C-terminus.

[0123] The amplified cDNA was cloned into expression cassette pCR-TOPO (plasmid from Invitrogen, Carlsbad, CA) following the vender's instruction and the sequences were verified by sequencing of both strands (Retrogen, San Diego, CA). The amplified cDNA sequence was identical to the sequence deposited in GenBank referenced above. The full-length Chk1 cDNA was constructed from these two overlapping cDNAs, ligating through the Clal restriction site at 734-739. This full-length cDNA was used as PCR template to generate cDNA fragments for expression or directly to generate the full-length Chk1 expression vector. All the PCR products were cloned into pCR-TOPO for sequencing. Constructs were made for the expression of full-length Chk1 and various lengths of kinase domain with or without 6xHis-tag.

Table 1

	PCR Primers*	
Primer	• Sequence	SEQ ID NO.
chk6w	GAG CTC AGT ACC ATC TAT CTT TTT TGA TGT CTG G	3
KH28	GAG CTC AGT TGG TGG TGG TGG TGT CCA CTG GGA GAC TCT	4
9	GAC AC	
K289	GAG CTC ATC CAC TGG GAG ACT CTG ACA C	5
Chk11	CCA TGG AGC TCA AGA AAG GGG CAA AAA GG	6
K210	GAG CTC ATT GGT CCC ATG GCA ATT CTC C	7
KH21	GAG CTC AGT GGT GGT GGT GGT GGT CCC ATG GCA ATT	8
0	стс с	
K248	GAG CTC ACT CAA CTA AGA TTT TAT GCA GCA G	9.
KH24	GAG CTC AGT GGT GGT GGT GGT GCT CAA CTA AGA TTT TAT	10
8	GCA GCA G	

3. Chk1 Antibodies

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[0124] Peptide NRVTEEAVAVKIVDMKRAVD (residues 28-47 of SEQ ID NO. 2) was selected for generating antibody against N-terminus of human Chk1. Peptide DDKILVDFRLSKGDGLE (residues 434-450 of SEQ ID NO. 2) was selected for generating antibody against C-terminus of human Chk1. Rabbit polyclonal antibodies were ordered through the Custom Antibody Production Services from Research Genetics, Inc. (Huntsville, AL). Both antibodies detected recombinant or endogenous human Chk1 as expected.

4. Fermentation

[0125] The overall scheme was follows. The 3' PCR primers were engineered to encode both untagged and tagged (with 6-histidine tag) proteins. The segment of cDNA for 1-289 was cloned into a pFastBac plasmid (obtained from Life Technologies) and an Ndel site was introduced. A recombinant baculovirus was generated using the Bacmid system (obtained from Life Technologies). The protein (KH289) was expressed in Hi-5 insect cells and purified by a combination of ion-exchange and affinity chromatography. The segments of cDNA for the full-length Chk1 (1-476AA) and the Chk1 kinase domain (1-265AA) were cloned into pAcSG2 plasmid and recombinant baculovirus was generated using BaculoGold viral DNA (obtained from Invitrogen) and a modified CellFectin transfection (obtained from Life Technologies) and plaque selection (obtained from Novagen) protocol. The expressed protein was purified using the chromatog-

raphy scheme described below. High salt concentration in buffers was found to be required to prevent precipitation of the purified proteins. The details of the protocol are discussed below.

Generation of Expression Plasmids

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[0126] Plasmid pFastBac-Nde was modified from the pFastBac1 vector (Life Technologies, Gaithersburg, MD) by in vitro site-directed mutagenesis using the Muta-Gene in vitro Mutagenesis Kit (Bio-Rad, Hercules, CA) following the supplier's instruction. Two nucleotides were substituted in pFastBac1 using the following oligonucleotide:

TGA ATA ATC CGG CAT ATG TAT AGG TTT TTT [SEQ ID NO. 14]

This created a unique Ndel site at the original translation start site for the polyhedrin protein.

[0127] The amplified cDNA fragments were digested with the restriction enzyme Stul and Sacl and cloned to plasmids pET28a (Novagen, Madison, WI), pAcSG2 (Pharmingen, San Diego, CA), or pFastBac-Nde. The pET28a vector was used for protein expression in *E.coli* and pAcSG2 and pFast-Bac-Nde were used for protein expression in insect cells. To clone the cDNA fragments encoding Chk1 kinase domain with amino acids 1-289 (construct KH289) into the pFastBac-Nde, the cDNA fragment was excised from the pCR-TOPO plasmid with restriction enzymes Stul and Sacl, ligated between the blunt-ended Ndel site and Sacl site. Plasmids with correct insertion were analyzed by restriction enzyme digestion. The full-length Chk1 and the kinase domain of residues 1-265 (KH265) with or without C-terminal 6xHis-tag were cloned into pAcSG2 using the restriction sites of Stul and Sacl. Expression vectors for kinase domain of residues 1-210 (KH210) and kinase domain of residues of 1-248 (KH248) were made in pFastBac-Nde.

[0128] Expression in E.coli was done following the instructions supplied with the pET28a vector. Proteins expressed in the form of full-length Chk1 or kinase domain of residues 1-265 or kinase domain of residues 1-289 were in the insoluble fraction when analyzed by ReadyPreps Protein Preparation Kit (Epicentre Technologies, Madison, WI).

Generation of Recombinant Viruses

[0129] The Bac-to-Bac system (Life Technologies) was used to generate recombinant baculovirus for expression of the C-terminally 6xHis-tagged Chk1 kinase domain (amino acids 1-289, KH289) as instructed. Recombinant viruses were confirmed by PCR for the presence of Chk1 cDNA insertion. Protein expression was confirmed by SDS-PAGE or Western blot with the Chk1 polyclonal antibodies. The expression of KH289 appeared to be the highest among all the constructs. High titer stocks of recombinant viruses were generated by 2 to 3 rounds of amplification using Sf21 insect cells.

[0130] Recombinant viruses for expression of the full-length Chk1 and kinase domain of residues 1-265 were generated by co-transfection of Sf21 cells with pAcSG2 vector and BaculoGold (PharMingen, San Diego, CA).

Expression in Insect Cells

[0131] The yield of active soluble protein obtained in the *E. coli* fermentation described above was impractical for large-scale experimentation. Therefore, an alternate fermentation system was developed. Insect cells Sf9 for viral amplification, and Hi-5 cells for protein production (both from Invitrogen, Carlsbad, CA, USA) were adapted to grow in insect medium contained 1% Fetal Bovine Serum (Life Technologies, Grand Island, NY, USA). Cells were propagated and maintained in suspension culture at 27°C in either Erlenmeyer shake flask (Corning # 430183) or in an upright roller bottle (Corning Inc., Corning, NY, USA # 25290-17000) with a loosened cap for aeration. The flasks were placed in a reciprocal refrigerated shaker (Innova 4343, New Brunswick Scientific, Edison, NJ, USA) at 120 rpm. The cell density was maintained at between 5 X 10⁵ to 2 X 10⁶ cells/ml by diluting the cultured cell suspension with a fresh pre-warmed (27°C) medium. The viability of insect cells was maintained at 98%. The viability of insect cells were determined by microscopic count of total stained cells by trypan blue versus the total cell number in a hemocytometer.

[0132] Sf9 insect cells were used for amplification for recombinant virus stock. The recombinant baculovirus from a single plaque was pick up by a pipette tip and added to Sf9 cells monolayer in T-25 flask (Becton Dickinson Labware, Franklin Lakes, NJ, USA) with 10 ml medium SF900II and 1% of Fetal bovine Serum (Life Technologies, Grand Island, NY, USA) and incubated at 27°C. After 6 days, the culture supernatant was used as first generation of virus stock (P1) for further amplification of P2 and P3 virus stocks to 2-3 L. For large scale amplification of the P2 and P3 virus stock, P1 or P2 virus stock was added to Sf9 cells at a cell density of 1 X 10⁶ cells/ml, the infection was carried out with Multiplicity Of Infection (MOI) of 0.1, cells were grown in suspension in 500ml of SF900II in 2 L roller bottle (Corning Inc., Corning, NY, USA) standing up right in a shaker incubator at 120 rpm at 27 ° C for 6 days. This process was repeated until 2-3 L viral stock (P3) were obtained. The titer of this virus stock was 1 to 5X10⁸p.f.u/mL. The viral titration was determined by the plaque assay method, with serial 10-fold dilution up to 10⁸ fold. The viral stock was stored at 10°C.

and used for large scale protein production within 2 months to avoid viral instability.

[0133] The Hi-5-insect cells (derived from Trichoplusia:ni-cells) which have been-adapted to grow in medium Ex-cell 401 (JRH Biosciences, Lenexa, KS, USA) with 1% Fetal Bovine Serium were used for protein production. The cells were grown in the upright roller bottle up to cell density at 2 X 10⁶ cells/ml; and were used as seed cells for bioreactor culture. The cells were grown in a 20 L stirred bioreactor with working volume at 18L (Applikon Inc., Foster City, CA, USA). Air flow rate was operated at about 10 ml per min per liter culture fluid. The air was fortified by pure oxygen in order to maintain the Dissolve Oxygen (DO₂) at 50% of air saturation. The agitation was maintained at 200 rpm throughout the cultivation. Cell density was started at about 5 X 10⁵ cells/ml and cells were infected when the density reached 2 x 10⁶ cells/ml. The MOI was 3 and the infection was carried out for 48 Hrs. After 48 hrs. of infection, the infected cells were harvested by centrifugation at 3,000 rpm for 10 min, at 4°C by a refrigerated centrifuge (model PR-7000M, IEC, Needham Heights, MA, USA). The cell pellets were collected and stored at -80°C.

5. Purification

6X-His tagged KH289

The basic purification scheme is depicted in Figure 4. Frozen cell pellets were thawed, suspended in ice-cold lysis buffer, and lyzed by microfluidizer (Microfluidics Corporation, Newton, MA). The lysis buffer contained 25 mM Tris-HCl, pH 8.0, 500 mM NaCl, 20 mM imidazole, and 14 mM __-mercaptoethanol. The lysate was centrifuged for 40 minutes at 40,000 rpm in a Ti45 rotor in Beckman L8-70M ultracentrifuge. The soluble fraction was flowed through a 150 mL Q-Sepharose FastFlow anion exchange column (Pharmacia, Piscataway, NJ), then loaded onto a 40ml Ni-NTA agarose column (Qiagen, Santa Clarita, CA). After extensive washes with the lysis buffer, the column was eluted with 240 ml of 20 mM to 300 mM imidazole gradient in the lysis buffer. Fractions containing the Chk1 kinase domain (KH289) were identified by SDS-PAGE and pooled. The pooled fractions were dialyzed in 25 mM Tris-HCl, pH 7.5, 500 mM NaCl, 0.5 mM EDTA, and 5mM DTT overnight. The dialyzed pool was diluted with 1.5 volumes of 25 mM Tris-HCl, pH 7.5, 20 mM MgCl₂, 8% glycerol, 5 mM DTT and loaded immediately onto a 40 ml ATP-Sepharose column. The column was eluted with 200 ml of 25 mM Tris-HCl, pH 7.5, 500 mM NaCl, 5 mM DTT, and 5% glycerol. Fractions containing KH289 were pooled and concentrated in a Millipore Stirred Cell under 60 psi N₂ and loaded onto a 320 ml HiPrep Sephacryl gel-filtration column and eluted with the same buffer. Pooled fractions were concentrated to 7-7.5 mg/ml for crystallography or ~3 mg/ml for HTS. Protein was flash-frozen in liquid N₂ and stored at -80°C.

[0135] Maintaining salt concentration around 500 mM NaCl including 5% glycerol was found to be crucial for preventing aggregation of Chk1 proteins during purification and storage without affecting the intended use.

6X-His tagged KH265 and KH476 Chk1

[0136] Essentially the same methods were used to purify the full-length Chk1 and the kinase domain of residues 1-265 expressed in insect cells. The expression protein levels as measured after the Ni-NTA chromatography or the final yields were much lower than that of the KH289 (full length sequence).

[0137] Gel-filtration HPLC has been used as a means of quality control. No significant difference was observed for samples stored at room temperature, 4°C, or -80°C for 4 days. The material eluted at a void volume that was less than 0.1%.

6. Crystallization, Crystallography and Three-Dimensional Analysis

[0138] The full length Chk1 protein (1-476 AA) had proven to be difficult to crystallize until the active kinase domain (1-289 AA) was identified. This active kinase was able to be expressed at the high concentration required for use in HTS and crystallography. The Chk1 data set was collected on MarlP345 under cryotemperature with stream freeze. The HB2-092 kinase domain preparation (1-289 AA) was first used. The initial data set at 2.35 () was obtained with overall Rsym of 4.6% and overall mosaicity for the data set is 1.2. Subsequent experiments with the HB2-101 (also a 1-289 clone) reached a 1.7 () resolution with mosaicity of 0.38 for the kinase domain using a crystal grown in refined conditions. Both the original and subsequent crystals have a space group P21 with one molecule per asymmetric unit. The results from the crystallographic analysis are shown in Table 2 below.

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Table-2:-Statistics	for the crystal	ographic analysis
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Crystal	Nat1	Nat2	AMP-PNP	Hg	Au
Internal merging and scaling				•	
Resolution (Å)	1.7	2.1	1.7	2.4	2.0
Reflections measured	162418	46947	107449	64881	125728
Unique reflection	35032	19145	35285	12821	22086
Completeness (%)	93.6 (88.3)	95.4(94.6)	94.1 (91.1)	95.4 (96.4)	97.5 (84.8)
Average I/o	29.9 (9.0)	15.47(4.38)	26.4 (12.5)	27.1 (11.6)	33.5(14.8)
R _{sym} ¹	3.2 (18.1)	5.0(23.3)	3.0 (10.0)	6.0 (13.2)	4.2 (11.8)
SIRSAS analysis					
Resolution (Å)				15-3.0	15-3.0
Rcullis ²		•		0.49	0.55
Phasing power ³ (SIR/SAS)				2.27/1.98	2.39/1.48
Figure of merit (combined)					0.764
Refinement statistics					
Resolution range (Å)	7-1.7	7-2.1	7-1.7		
Reflections used ⁴ (F>1oF)	30132	15804	31794		
Total nonhydrogen atoms	2372	2354	2460	•	
Rcryst ⁵ (%)	21.6	20.8	22.6	•	
Rfree ⁶ (%)	23.5	25.0	24.9	•	
rmsd from ideal bond length (Å)	0.005	0.006	0.010		
rmsd from ideal bond angle (°)	1.30	1.27	1.58		
Average B ($Å^2$; all atoms)	28.9	29.7	23.22		

Data for the outermost resolution shell are given in parentheses.

N _ N

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h i=1 h i=1

where I(h)i is the ith measurement of reflection h and I (h) is the mean value of the N equivalent reflections.

- 2 Rcullis = 11 FPH +/- FP 1 FH(calc) 1 / 1 FPH +/- FP 1 for all centric reflections.
- 3 Phasing power = r.m.s. (IFHI/E), where IFHI is the heavy-atom structure factor amplitude and E is the residual lack of closure.
- 4 Number of reflections used in working set.
- 5 Rcryst = 1 |Fobs| 1 |Fcalc| |/•|Fobs|, where summation is over data used in the refinement.
- 6 Riree is the same calculation including the 10% of data excluded from all refinements.

Crystals were grown at 13°C using a hanging-drop vapor-diffusion method. Two crystallization conditions produced the exact same form of crystals. The Nat1 crystal was obtained by mixing equal volume of protein solution (7to 7.5 mg/ml protein) and reservoir solution of 13% PEG 8000 (w/v), 0.115 M (NH₄) $_2$ SO $_4$ 0.1 M NaCacodylate (pH 6.8), 2% glycerol. The Nat2 crystal was crystallized using reservoir solution of 12% PEG8000 (w/v), 15% isopropanol, 0.1 M Hepes (pH 7.5). The crystals belong to the space group P2₁ and have unit cell dimensions $a = 45.2\text{\AA}$, $b = 65.7\text{\AA}$, $c = 65.7\text{\AA}$ 58.1Å, d = 93.9°. The crystals contained one molecule per asymmetric unit and are 53% solvent by volume. The crystals of binary complex with AMP-PNP were obtained by co-crystallization first under the same crystallization condition as Nat1 crystal in the presence of 1.25 mM AMP-PNP and 2.5 mM MgCl₂, then the resulting crystals were soaked in mother liquor containing 5 mM MgCl₂ and 20 mM AMP-PNP for two days. The co-crystals had the identical space group (P2₁) and cell dimensions as the native crystals. All diffraction data were collected at -170°C. Crystals were introduced into cryoprotectant solution containing its reservoir solution and 20% glycerol. For AMP-PNP co-crystal, additional 10 mM MgCl₂ and AMP-PNP were included in cryoprotectant solution. Crystals were then flash frozen in a stream of nitrogen gas -170°C. All data collection was carried out with home source using CuK γ-radiation produced by a Rigalu rotation anode FR5 X-ray generator equipped with focusing mirrors and measured with a Mar 345 image-plate detector. All data were processed with the Denzo/HKL package (Otwinowski, Z., *Oscillation Data Reduction Program*, Proceedings of the CCP4 Study Weekend: Data Collection and Processing, pp. 56-62, compiled by: L. Sawyer, et al., SERC Daresbury Laboratory, England (January 29-30, 1993)).

¹ $R_{sym} = \cdot \cdot \cdot I(h) - I(h)_i V \cdot \cdot I(h)_i * 100,$

Initial apoenzyme structure determination using Nat1 crystal data was carried out by molecular replacement (MR) using modified Cdk2 structure (omitted loop regions) (Russo, AA et al., Nature 382(6589):325-31 (Jul 25, 1996)) as a search model. Rotation and translation functions using the AMoRe software (Navaza J, Acta Crystallographic, 50(2): Section A (March, 1994)) revealed a solution using Nat1 data from 10 to 4 Å. The MR model was refined by simulated annealing (X-plor). However, after successive rounds of rebuilding and refinement, 2Fo-Fc and Fo-Fc electron density maps were poorly defined at the loop regions which were omitted from the initial model. To obtain additional phase information, multiple isomorphous replacement was carried out with two heavy metal derivatives: 0.5 mM HgCl₂ (soaked for 15 hrs) and 5 mM Kau (CN)2 (soaked for 17 hrs). Five Hg sites and five Au sites were identified by difference Fourier synthesis using phases generated from the MR partial model and were consistent with both isomorphous and anomalous difference Patterson maps. The positional and thermal parameters and relative occupancies for the heavy atom sites were refined using SIR data at 3 Å and anomalous data at 3.5 Å by program PHASES (Furey, W et al. *Phases: a Package of Computer Programs Designed to Compute Phase Angles for Diffraction Data from Macromolecular Crystals*, American Crystallographic Association, Series 2, 18:73 (1990)). Sixteen cycles of solvent flattening were then carried out using phases calculated from refined Hg and Au positions. The resultant electron density maps showed a good backbone density and well-defined side chains for most part of the protein. Model building utilized the program FRODO (Jones, T.A., J Appl Cryst, 11: 268-272 (1978)). The missing loop regions were incorporated into the model using both MIR maps and model phased 2Fo-Fc maps. Further refinement in XPLOR (Brünger, A.T. et al., X-PLOR Version 3.1: A System for X-ray Crystallography and NMR", Yale University Press, (1992)) and then CNS (Brünger, A.T. et al, Crystallography & NMR System, Acta Cryst., D54: 905-921 (1998)) were continued with both conjugate gradient minimization and simulated annealing, then followed by manually rebuilding.

[0141] Refinement of Nat2 structure was carried out by using refined Nat1 model but omitting residues 153-170 as well as SO₄. Fo-Fc maps showed well defined densities for the omitting region and its conformation is exactly same as that in Nat1.

[0142] Refinement of the binary complex with AMP-PNP was proceeded with refining the position of the refined apo-enzyme model (Nat1) as rigid body against the complex data using CNS program. Fo-Fc maps with __A*(Read, R.J., Acta Cryst., A42: 140-149 (1986)) weighting showed clear density for the adenine and ribose components of AMP-PNP. The conformation of residues forming the binding pocket was checked in simulated annealing omit maps before including the adenine and ribose components of AMP-PNP.

The apo-enzyme model (Nat1) included all atoms for residues 2 to 44 and 48 to 276, 183 ordered solvent molecules and one SO₄ molecule. The refined Nat2 structure contained the same number of residues and solvent molecules but the SO₄ molecule was not present. The refined AMP-PNP complex contained the same number of residues as apo-structures, with 150 ordered solvent molecules and one SO₄ molecule. The triphosphate moiety of AMP-PNP was disordered and no Mg²⁺ ions were visible. The final model had all residues in "most favored" or "additional allowed" regions of the Ramachandran plot according to PROCHECK (Laskowski RA et al., *J. Appl. Cryst.*, 26: 283-291 (1993)), with no residues in "generously allowed" or "disallowed" REGIONS, indicating the well refined nature of the identified crystal structure. The terms "generously allowed" and "disallowed" are descriptions of the configuration of Phi and Psi angles of the protein structure. A well refined protein structure should not place these angles in the unpreferred or non-naturally occurring configurations.

7. The Overall Kinase Structure

[0144] The crystal structures of the kinase domain of human Chk1 and its binary complex with an ATP analog, AMP-PNP, have been determined to 1.7 Å resolution. Both structures contain the kinase core domain (residues 2-267) and residues in the linker region that connects the N-terminal kinase domain with the C-terminal region of Chk1. The crystallographic analysis is summarized in Table 2. The Chk1 crystal coordinates for the apoenzyme (isolated active Chk1) and the binary complex (Chk1 complexed with AMP-PNP, an ATP analog) are shown in Figures 11A and 11B, respectively. The coordinates of the fixed water molecules are also included therein.

The kinase domain of human Chk1 has a canonical kinase two-lobe fold, with the ATP binding cleft between the two lobes (Figure 5, structure model). The smaller N-terminal lobe contains one helix (α C) and 5 β -strands (β 1 to β 5) that form a curved anti-parallel β -sheet. The larger C-terminal lobe contains a cluster of 7 helices (α D to α I), packed against 6 β strands (β 6 to β 11) which border the cleft. One β strand (β 6') comprises the hinge region connecting the two lobes. In both apo-enzyme and binary structures, the ATP binding site, catalytic residues, and the activation loop are well ordered. Comparison with crystal structures of other kinases indicates that the Chk1 kinase domain is closely related to PhK (Lowe, ED et al., *EMBO J*, 16(22):6646-58 (Nov 17, 1997)) (See Figure 1A, 1B). The N-terminal lobe (Residues 2-90) superimposes with an r.m.s. derivation for C α atoms of 1.1 Å, while the C-terminal lobe (Residues 91-276) superimposes with an r.m.s. derivation for C α atoms of 0.9 Å. In the C-terminal lobe, major differences are found in helix α G, and the connecting loop between α G and α H. These are not included in the superposition. The Chk1 apoenzyme adopts a more open conformation compared to PhK. The N-terminal lobe of Chk1 is rotated ~15° relative

to the ternary complex of PhK with its substrates. Comparison of the AMP-PNP bound Chk1 binary complex with the apoenzyme structure shows no conformational change. A high degree of sequence homology for Chk1 kinase domains of different species (Figure 2) suggests that there is an overall structural conservation of the kinase domain. Residues that are not modeled in the current structures are not conserved in Chk1. For example, there is a six-residue insertion in the loop connecting $\beta 3$ and αC in S. pombe Chk1.

The two lobes are held together by an extensive hydrogen-bond network at the lobe interface which involves the loop linking αC and $\beta 4$ of the N-terminal lobe, $\beta 6$ ' of the hinge region, and $\beta 7$ and $\beta 8$ of the C-terminal lobe. This network extends from the back of the protein to the front opening of the ATP binding cleft. Residues involved in this network also form part of the pocket that interacts with the adenine moiety of AMP-PNP. Strand $\beta 8$ immediately precedes the kinase conserved DFG motif, in which Asp148 is important for the alignment of the phosphate groups of ATP. The only reported mutation in the Chk1 kinase domain is at the lobe interface. Replacement of the conserved Glu85 by Asp leads to a temperature-sensitive phenotype in fission yeast in which the mutant maintains cell cycle arrest after UV irradiation but impairs the DNA replication checkpoint at nonpermissive temperature (Francesconi, S et al., *EMBO J*, 16(6):1332-41 (Mar 17, 1997)). The side chain of Glu85 at the end of strand $\beta 5$ forms hydrogen bonds with the side chain of conserved Lys145 from strand $\beta 8$ as well as with the main chain amide of conserved Lys69 that precedes strand $\beta 4$. These interactions, together with the extensive hydrogen-bond network at the lobe interface, appear to play an important role in maintaining the correct disposition of the N-terminal lobe and the DFG loop during lobe movement. The Glu to Asp mutation, while maintaining similar charge, would not be long enough to form those hydrogen bonds provided by Glu85, thereby weakening lobe interactions and rendering the mutant protein less stable at higher temperature.

[0147] Most of the invariant residues of Chk1 proteins are located in the C-terminal lobe. Many of them are also conserved among Ser/Thr kinases and are involved in stabilizing the catalytically active kinase conformation and in binding ATP. The positions of several invariant motifs of Chk1 proteins are noteworthy. Compared with other Ser/Thr kinases, the IEPDIG motif (residues 96-101) shortens αD to a one-turn helix, since Pro98 initiates a tight turn between αD and αE. This turn interacts with the C-terminus of helix αF through a backbone hydrogen bond between Asp99 and the invariant Gly204. In this turn, Glu97 forms backbone hydrogen bonds with Ile100 and Gly101. The unique conformation of this motif appears to be important for peptide substrate interaction, since the side chains of Ile96 and Pro98 form part of a hydrophobic pocket that interact with the peptide substrate as discussed below. Helix αE contains a conserved motif of AQXFFXQL (residues 107-114; SEQ ID NO: 24), with the hydrophobic residues buried inside the C-terminal lobe. The side chain of Gln108 projects towards the linker region that follows the kinases core domain and forms hydrogen bonds directly or through a water molecule to backbone atoms of Lys267, Leu269 and Lys270. Although Chk1 sequences diverge in this linker region, these backbone interactions with Gln 108 could still be conserved, holding the linker against the N-terminus of αE . Helix αG is positioned differently compared with αG of PhK. Two sets of invariant PW residues (207 and 208, 230 and 231) flanking αG, although separated by 21 residues, are in van der Waals contact and connected to the hydrophobic core of the C-terminal lobe. This stabilizes the surface for peptide substrate interaction.

Activation and Catalytic Loops

Interesting features of the Chk1 kinase domain include interactions that stabilize the activation loop. The structure of the activation loop determines the alignment of residues contacting ATP and performing catalysis in protein kinases. Interacting with the catalytic loop, the activation loop orients the catalytic Asp; interacting with the N-terminal lobe, the activation loop closes the N and C terminal lobes and aligns residues that interact with the phosphates of ATP. The activation loop is defined as the region between the conserved motifs of DFG and APE corresponding to residues 148 to 177 of Chk1. Conformational changes in the activation loop serve as a major regulatory mechanism for kinase activity. In the human Chk1 structures, the activation loop is folded in a conformation similar to those found in structures of active kinases, consistent with the observation that the Chk1 kinase domain is constitutively active. This active conformation is stabilized by special features of Chk1 secondary structures and their side chain interactions (Figures 3 and 5, homology model and crystal structure).

[0149] The N-terminus of the activation loop interacts with the catalytic loop through the interaction of $\beta6$ and $\beta9$. Immediately following $\beta9$, $\beta10$ interacts with $\beta11$ to form a two-stranded β -loop with a turn at Asn159. This β -loop is packed against the N-terminus of the catalytic loop and positions the highly conserved Arg156 and Glu161. The side chain of Arg156 interacts with the carbonyl of the invariant His122 at the end of αE . Through the invariant Asp190, the side chain of His122 is connected to the amide of Arg129, adjacent to the catalytic residue Asp130. The carboxyl of Glu161 forms a hydrogen bond with the imidazole of His185 that precedes αE . These interactions anchor this end of the activation loop to the core of the C-terminal lobe. The center of the activation loop interacts with the rest of C-terminal lobe through two backbone hydrogen bonds between Leu164 and Phe184. The activation loop ends at its C-terminal with a turn which is supported by αEE . In human Chk1, αEE is anchored at two positions to the core of the C-terminal

lobe through two ion-pairs, one is the invariant kinase ion-pair between Glu177 and Arg253, another is between Lys180 and Glu248 which is unique to Chk1. This extra ion-pair constrains the movement of αΕΕ, and in turn the movement of the C-terminal end of the activation loop. The pair of Lys180 and Glu248 is only conserved in vertebrate Chk1, suggesting potential flexibility of αΕΕ and the activation loop of Chk1 in lower organisms such as S. pombe.

[0150] Crystal structures of kinases indicate that the conformation of the activation loop is influenced by its negative charge which neutralizes a cluster of positively charged residues, although the ionic interaction may not be absolutely required as in the case of mammalian casein kinase I. The negative charge is provided by phosphate through phosphorylation, carboxyl group of Glu, or solvent ions. In Chk1, the positively charged cluster of Arg129, Arg162, Lys166, and Lys54 is present, but no phosphorylation is observed. In both the apoenzyme and binary complex structures determined to 1.7 Å, a sulfate ion was close to the phosphate position of the phosphothreonine (Thr197) in PKA. This sulfate ion interacts with Arg129, Arg162, and Thr153. Sulfate is present in the crystallization solution and could contribute to the stability of the positively charged cluster and the activation loop. To clarify the role of this sulfate ion and to better understand the interactions that stabilize the activation loop, crystals were produced under sulfate-free condition and determined the structure to 2.1 Å (Table 2). This 2.1 Å structure is referred as Nat2 structure, whereas the 1.7 Å apoenzyme structure is referred as Nat1 structure. In Nat2 structure, no sulfate ion is present.

[0151] Superimposition of Nat1 and Nat2 structures revealed similar conformations for the corresponding activation loops except for the side chain of Arg162 which turns toward the solvent in Nat2 structure. The side chain of Arg162 is flexible in both structures as indicated by its high temperature factors. Arg162 is an invariant residue of Chk1 and its function is not readily apparent from the structure. In both the Nat1 and Nat2 structures, the side chain of Arg129 forms hydrogen bonds to three main chain carbonyl oxygens (Leu151, Ala152, and Lys166) directly or via water molecules. The positive charge of Arg129 could be neutralized by the thiol group of Cys168 which is in the vicinity of side chains of Lys166 and Arg129. In this basic environment, this thiol could become a thiolate ion. Cys168 is invariant in Chk1 and is conserved in many kinases such as PKA and PhK. Our results rule out the role of sulfate ion in stabilization of the activation loop of Chk1. Instead, the activation loop and the catalytic loop are stabilized by its unique secondary structures and their extensive side chain interactions.

[0152] A difference between Chk1 and other kinases is the permuted positions of Lys166 and Thr153 (Figure:2). Lys166 occupies the equivalent position as Giu182 of PhK and the phosphorylated Thr197 of PKA, whereas Thr153 is equivalent to Lys189 of PKA. The side chain of Thr153 forms a hydrogen bond with the side chain of Lys54 located in helix αC. Thr153 is conserved in Chk1 (Thr or Ser) and is a candidate for phosphorylation in the activation loop. The permuted position, however, makes phosphorylation of Thr153 unlikely. The activation loop is already in an active conformation in Chk1 and phosphorylation would be unnecessary. Lys54 is conserved in all but S. pombe Chk1 and adjacent to Glu55 which forms the invariant ion-pair with Lys38 in active kinases. The interaction between Thr153 and Lys54, therefore, appears to play a similar role to the interaction between His87 and the phosphate of Thr197 of PKA. The side chain of Lys166 points to Cys168 and its position appears to play a role in determining the substrate specificity as discussed below. In S. pombe Chk1, the residue that corresponds with Lys166 is Ser, suggesting potential regulation of the activity of S. pombe Chk1 through phosphorylation. Concomitantly, the activation loop of S. pombe Chk1 appears to be more flexible since its substitutions would disrupt some of the interactions that stabilize the activation loop.

Catalytic Residues and AMP-PNP Binding

[0153] The glycine-rich loop that anchors the phosphate groups of ATP in kinases is poorly ordered in Chk1, as evidenced by the high B factors in this region for both apoenzyme structures and AMP-PNP bound binary complex structure. Residues 18-21 at the apex of the loop between β 1 and β 2 are flexible with poor electron density. These residues are highly conserved in kinases and anchor the β -phosphate of ATP in ATP-bound kinase structures. The flexibility of this loop could play a role in regulating Chk1 kinase activity, indeed, Tyr20 present in higher organisms corresponds structurally to Tyr15 of Cdc2 which following phosphorylation inhibits Cdc2 activity (Coleman TR, et al., *Curr Opin Cell Bio*, 6(6):877-82 (Dec. 1994); Russo, AA et al., *Nature*, (1996), supra).

[0154] One striking feature among the active ternary complexes such as PKA and PhK is the close similarity of the active site residue conformation, their interactions with the ATP and coordination of the metal ions. The binary complexes that have been solved show no such conservation (Knighton DR, et al., *J Mol Biol*, 220(2):217-20 (Jul 20, 1991); Bossemeyer, D et al., *EMBO J*, 12(3): 849-59 (Mar 1993); Zheng J, et al., *Protein Sci*, 2(10):1559-73 (Oct 1993); Owen DJ, et al., *Structure*, 3(5):467-82 (May 15, 1995); Lowe, et al., *EMBO J*, (Nov 17, 1997), supra.). Many of the active site residues in the Chk1 structure have interactions quite similar to those in ternary complexes of Phk and PKA (Figure 4A, 4B). In the N-terminal lobe, the invariant ion pair of active kinases is present between Lys38 and Glu55; the corresponding Lys in PhK and PKA interacts with α and β phosphates of ATP. Helix αC is firmly attached to the rest of N-terminal lobe through hydrophobic interactions and is in an active position relative to the rest of the N-terminal lobe. It also interacts with the DFG loop in the C-terminal lobe, the side chain of Glu55 from αC rests above Gly150. The relative side chain positions of Lys38, Glu55, and Asp148 are similar to those for the corresponding residues in the ternary com-

plexes of PKA and PhK. These residues in PKA and PhK, together with the glycine-rich loop, coordinate a Mg2+ and anchor the lpha and eta phosphates of ATP. In the C-terminal lobe, the conformation of the catalytic loop (residues 130-135) of Chk1 is nearly identical to that in PhK with the side chains of Asp130, Lys132, and Asn135 in Chk1 nearly superimposable to the corresponding residues Asp149, Lys151, and Asn 154 in PhK in which Lys151 binds to the γ -phosphate of AMP-PNP and Asn154 chelates another Mg2+ that binds to the β and γ phosphates of AMP-PNP. Thr170 is conserved in all serine/threonine protein kinases and appears to determine the specificity of Ser/Thr verses Tyr as phospho-acceptor. Thr170 forms hydrogen bonds with Asp130 and Lys132 analogous to Thr186 in PhK and these interactions are needed for the positioning the carbonyl of the catalytic residue Asp130. The residues of Chk1, however, are far apart from those in the N-terminal lobe and the DFG loop due to the somewhat open lobe conformation (Figure 6). The DFG loop is positioned higher than its counter parts in PKA and PhK. Lys38Ns is 10 Å away from Asp130082, compared with 8.2 Å in Phk and 7.8 Å in PKA. Asp148081 is 6 Å away from Asp130082, compared with 3.8 Å in PhK and 4.8 Å in PKA. In Chk1, one water molecule is located between Asp148 and Asp130 and is hydrogen bonded to Asp1300δ2 as well as Asn1350δ1. The side chain of Asn135 is over 1 Å farther away from Asp148 relative to the active conformation in PhK. The residues that are necessary for ATP phosphate binding and catalysis are clustered in two separate parts, although they maintain their local interactions. The lack of electron density of the triphosphate moiety of AMP-PNP in the binary complex of Chk1 probably results from misalignment of these residues as well as flexibility in the glycine-rich loop.

The adenine and ribose moieties are clearly defined in our current model. As in all the structures of kinases with ATP, the adenine base is almost completely buried in a hydrophobic pocket between the two lobes, and hydrogen bonds are formed between N6 of adenine and the main chain carbonyl of Glu85, and between N1 and amide of Cys87. As in PhK, Chk1 N7 interacts with the side chain of Ser147 via a water molecule in Chk1. However, the ribose ring adopts a C2'-endo conformation similar to that in the inactive form of Cdk2 (PDB ID code 1HCK, (De Bondt HL, et al., Nature, 363(6430):595-602 (Jun 17 1993); Schulze-Gahmen U et al., J Med Chem, 39(23):4540-6 (Nov 8, 1996)), with the O2' hydrogen-bonding to Glu91, and O3' hydrogen bonding to the carbonyl of Leu15 in the glycine-rich loop. In comparison, the ribose rings have C3'-endo puckering in the active ternary complexes of PKA and PhK:

Substrate Specificity and Interactions That Stabilize the Closed Conformation

[0156] The structured activation loop of Chk1 provided an opportunity to explore the basis of peptide substrate specificity. The close resemblance of Chk1 with PhK and the available structures of PhK with and without peptide substrate enable us to model the interactions of peptide substrate with Chk1. The interaction of kinases with their peptide substrates has been analyzed for three kinases, PKA with an inhibitor peptide of PKI (PDB code 1ATP, (Knighton DR. *J Mol Biol*, (Jul 20, 1991), supra.), PhK with MC-peptide (PDB code 2PHK, (Lowe, et al., *EMBO J*, (Nov 17, 1997), supra.), and insulin receptor tyrosine kinase with a peptide substrate (PDB code 1IR3, (Hubbard SR, *EMBO J*, 16(18):5572-81(Sep 15, 1997)). In all three tertiary complex structures, the backbones of peptide substrates around the phosphate acceptor residues adopt extended conformation and interact mainly with the C-terminal lobes.

[0157] The known Chk1 kinase substrate is the Cdc25C protein phosphatase. Several phosphate acceptor Ser residues have been identified in the Cdc25C protein sequence. Consensus features can be derived from sequences surrounding the phosphate acceptor Ser (position P): The N-terminal P-3 position is a conserved-Arg, P-5 positions prefers bulky hydrophobic residues, and P-2 is Ser or Thr. Phosphorylation of Ser216 of human Cdc25C is required for DNA damage induced G2 arrest and Ser216 is phosphorylated by Chk1 in vitro (Peng et al., *Science* (1997), <u>supra.</u>); Sanchez et al., *Science* (1997), <u>supra.</u>). Therefore, the peptide LYRSPSMPE spanning residues 211-219 of human Cdc25C was used to model the interaction of peptide substrate with Chk1, based on the ternary complex of PhK with MC-peptide.

[0158] The modeled Cdc25C peptide easily fits into a groove on the C-terminal lobe of Chk1, following a path very similar to that of the MC-peptide bound to PhK (Figure 7). The Oγ atom of Ser(P), the presumed nucleophile in the phosphate transfer reaction, is very close to an ordered water molecule in Chk1 structures. This water molecule hydrogen bonds to both the Asp130Oδ2 and Lys132Nε. Superposition of Chk1 and PhK shows that this water molecule would be 3.4 Å from the γ-phosphorus atom of the AMP-PNP in PhK. The position of this water molecule probably indicates the approximate location of the seryl hydroxyl during catalysis.

[0159] The hydrophobic side chain of Leu(P-5) fits into the hydrophobic pocket formed by Phe93, Ile96, Pro98, and Leu206. All of these residues except Leu206 are invariant in Chk1 proteins. The side chain of Arg(P-3) points towards Glu91 of Chk1. However, in its extended conformation, the guanidinium group of this Arg can only make a hydrogen bond (3 Å) with the carboxyl of Glu91. In both PKA and PhK, the guanidinium of Arg(P-3) forms a salt bridge (2.5 Å) with the carboxyl of the corresponding Glu residues. As discussed below, ionic interaction of Arg and Glu91 could be established after lobe closure.

[0160] The side chain of Ser(P-2) could make a hydrogen bond to the backbone carbonyl oxygen of Pro(P-1). In PhK, Gln(P-2) of the MC-peptide interacts with Ser188. This interaction is not available to Chk1 since it has an invariant

Pro172 in the corresponding position of Ser188 in PhK. Pro172, then, may contribute to the specificity of Chk1 for Ser or Thr at P-2 position and the internal hydrogen bond provided by Ser or Thr at P-2 position may play a role in maintaining the conformation of the substrate backbone at its N-terminus.

[0161] The hydrophobic side chain of Met(P+1) projects into a hydrophobic pocket formed by residues of Leu171, Val174, Leu178, Leu179, and Met167. The P+2 position can only accommodate a small side chain or a turn due to the unique position of Lys166. Lys166 is conserved among vertebrate Chk1 proteins. Correspondingly, Pro is found at the P+2 position of the Cdc25 substrates. Pro(P+2) creates a consensus 14-3-3 binding site once the Ser(P) is phosphorylated. The Lys166 of human Chk1 is a Ser residue in S. pombe Chk1. The side chain of S. pombe Chk1 could be phosphorylated and point to the position corresponding to the sulfate ion in human Chk1 structure. Correspondingly, bulky side chains are present at the P+2 position of the substrates of S. pombe Chk1.

[0162] Phosphorylation of Cdc25C by Chk1 is very specific such that the Ser(P-2) is not phosphorylated. This is important for Cdc25C regulation since phosphorylation at the P-2 position would destroy the 14-3-3 binding site. Our model clearly indicates determinants for Chk1 substrate specificity: hydrophobic interaction through the P-5 and P+1, ionic interaction through P-3, Ser/Thr at P-2, and small amino acid side chains at the P+2 position.

[0163] Although the recombinant Chk1 kinase domain is active when assayed in solution, the structure reveals that it is not in a closed catalytically active conformation in either the apoenzyme or the binary crystal structure. This result suggests that the apoenzyme and the ATP bound binary complex favor the open conformation. Lobe movement is common in kinase domains and catalysis requires a closed conformation (Cox S, et al., *Curr Opin Struct Biol*, 4(6):893-901(Dec, 1994); Gangal M, et al., *Biochemistry*, 37(39):13728-35 (Sep 29, 1998)). Interactions that stabilize the closed active conformation have not been addressed in detail in previous reports. Our model suggests that a key interaction in Chk1 is the ion-pair between Glu91 with Arg(P-3) of peptide substrate.

[0164] Superposition of Chk1 and PhK structures indicates that lobe closure of Chk1 can be achieved by a simple rotation of the N-terminal lobe by ~15 degree around residue Glu91. This rotation would place Glu91 closer to Arg(P-3) and establish an ion-pair between the carboxylate group of Glu91 and the guanidinium group of the Arg(P-3). Lobe closure could also change the ribose conformation of AMP-PNP to a C3'-endo conformation from the C2'-endo conformation in the binary complex. The catalytically active kinase ternary complex structures reported to date have their respective ribose rings puckered in a C3'-endo conformation. For Chk1, when the ribose is modeled in a C3'-endo conformation, two hydrogen bonds can form between the carboxyl group of Glu91 and the O2' and O3' of the ribose. In comparison, the binary complex of Chk1 with AMP-PNP has only one hydrogen bond between Glu91 and the ribose. The Chk1 kinase domain in solution likely shifts dynamically ("breathes") between the open and closed conformation. The current Chk1 structures have open conformations and have revealed that the ATP binding cleft is accessible to solution. In the closed conformation, residues for phosphate binding and catalysis come together and align the phosphate for transfer. The additional interaction of Glu91 with Arg(P-3) of peptide substrate and with the ribose of ATP would shift the equilibrium to the closed active conformation. Therefore, peptide substrates gain specificity partly through their ability to stabilize the closed catalytically active conformation of Chk1.

8. Regulation of Chk1 Kinase Activity

Phosphorylation of the Chk1 substrate, Cdc25, and the resulting cell cycle arrest has been correlated with the activation of Chk1 after DNA damage. Whether phosphorylation of Chk1 regulates its kinase activity is unclear. The structure of human Chk1 suggests that its activity is not regulated through phosphorylation of the activation loop. Instead, the activation loop of Chk1 appears to be anchored by extensive interactions through rigid secondary structures and their side chains. Interestingly, phosphorylation of the activation loop could occur in S. pombe Chk1 which has a Ser substitution at the position of Lys166. Whether Chk1 is regulated differently in S. pombe and mammals requires the identification of residues that are phosphorylated after DNA damage.

[0166] The structure of the Chk1 kinase domain and its binary complex with AMP-PNP provide insight into its activation mechanism. First, the structures reveal an unique arrangement of the residues for phosphate binding and catalysis. Specifically, the residues for α and β phosphate binding are separated from those for γ phosphate binding and catalysis. Alignment of these residues is achieved in a closed conformation which is stabilized by peptide substrate. Our model predicts low ATPase activity of Chk1 and favors an ordered kinetic mechanism in which ATP binding precedes the peptide substrate binding. Secondly, the structures exclude a role for the activation loop of human Chk1 in regulating the kinase domain conformation. The activation loop is most likely maintained by rigid secondary structures and the extensive interactions of their side chains. However, a possibility of different regulatory mechanism exists for S. pombe Chk1, which may reflect their different cell cycle processes and different DNA damage repair mechanisms. In addition, the interactions that stabilize the active kinase conformation have been identified. The presence of Glu in many kinase hinge regions and Arg at P-3 position of their substrates suggests a general role for this interaction in maintaining the closed conformation for Ser/Thr kinases. Interactions that determine the peptide substrate specificity suggest a consensus sequence that is useful to identify potential Chk1 substrate. Finally, Chk1 kinase domain structure provides a guide

for its future characterization as well as design of specific inhibitors that could abrogate checkpoint control for cancer therapy.

9. Enzymatic Activity of Chk1

The enzymatic activity of a kinase is measured by its ability to catalyze the transfer of a phosphate residue from a nucleoside triphosphate to an amino acid side chain in a selected protein target. The conversion of ATP to ADP generally accompanies the catalytic reaction. Herein, a synthetic substrate peptide, Syntide-2, having amino acid sequence PLARTLSVAGLPGKK (SEQ ID NO. 11) was utilized. The production of ADP from ATP that accompanies phosphoryl transfer to the substrate was coupled to oxidation of NADH using phosphoenolpyruvate (PEP) through the actions of pyruvate kinase (PK) and lactic dehydrogenase (LDH). The oxidation of NADH was monitored by following the decrease of absorbance at 340 nm (e340=6.22 cm-1 mM-1) using a HP8452 spectrophotometer. Typical reaction solutions contained: 4 mM PEP, 0.15 mM NADH, 28 units of LDH/mL, 16 units of PK/mL, 3 mM DTT, 0. 125 mM Syntide-2, 0.15 mM ATP and 25 mM MgCl₂ in 50 mM TRIS pH 7.5; 400 mM NaCl. Assays were initiated with 10 nM of kinase domain of Chk1, KH289. K_i values were determined by measuring initial enzyme activity in the presence of varying concentrations of inhibitors. The data were analyzed using Enzyme Kinetic and Kaleidagraph software.

[0168] The table below (Table 3) compares three different preparations of Chk1. The first preparation is the full length form, which comprises amino acids 1-476 of SEQ ID NO. 2. The next preparation contains proteolytically cleaved fragments, a mixture of Chk1 protein fragments obtained from the full-length protein during fermentation. The exact enzymes involved and cleavage site generated for these fragments is unknown. However, analysis of the fragments indicated that one of them is similar in size to the 1-289. The third preparation is the kinase domain of amino acids 1-289 of SEQ ID NO. 2 (KH289) As mentioned above, the assay used detects the ADP product by coupling through the enzymatic actions of pyruvate kinase and lactate dehydrogenase.

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Table 3

Prep No.	Prep	Concentration	Rate/min	Activity	Ki
HA2-013	Full Length Chk1	75nM	0.0190	1 (control)	48 ± 1nM
HA2-022	Proteolytically cleaved Chk1	2nM	0.0208	+38 fold	37 ± 5 nM
HB2-061	Kinase Domain Chk1 (1-289)	7.3nM	0.0200	+10 fold	68 ± 12 nM

[0169] Additional activity comparison experiments were performed using new preparations of full length Chk1, proteolitically cleaved Chk1, and kinase domain Chk1. The preparation conditions were as described above. Once again, the cleaved preparation was 38 fold more active than the non-cleaved preparation.

10. High Throughput Screens

[0170] T

The following substrates were tested for peptide content and activity:

Table 4

Peptide Substrates												
		Activity	Peptide									
Syntide 2	PLARTLSVAGLPGKK (SEQ ID NO. 11)	100%	75%									
Syntide 3	KAGAG-PLARTLSVAGLPG-Biotin-K (SEQ ID NO. 12)	67%	50%									
Syntide 4	Ac - PLARTLSVAGLPG-AGAGAGAK (SEQ ID NO. 13)	72%	45%.									
Syntide 5	PLARTLS (PO3) VAGLPGKK (SEQ ID NO. 15)	··NT	42%									
Syntide 6	PLARTLS (PO ₃) VGALPGK (Biotin) (SEQ ID NO. 16)	NT	77%									

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[0171] As described in detail below, an aspect of the invention involves a nonradioactive ELISA based assay suitable for high throughput screening (HTS). The development of the ELISA based CHK1 kinase HTS assay was first initiated with a monoclonal anti-phosphoserine antibody called Clone PSR-45, supplied by Sigma. New Chk peptide

substrates, analogues of Syntide2, were synthesized to validate this assay. These peptides are listed in Table 4. Biotin-Syntide-2 (SEQ ID NO. 12), and N-terminus acetylated Syntide-2 (SEQ ID NO. 13) and the expected peptide products after CHK phosphorylation, serine phosphorylated Syntide 2 (SEQ ID NO. 15), and serine phospholylated biotin-Syntide 2 (SEQ ID NO. 16) were synthesized for assay development. Although the assay worked well in solution with these peptides, it did not work when the peptide (serine phosphorylated Syntide 2.— SEQ ID NO. 15) was immobilized on DNA BIND (Costar) 96 well plates. This antibody also did not work well when the biotin-labeled peptide was immobilized using Neutravidin coated 96 well plates (Pierce). To circumvent these issues, a polyclonal antibody specifically directed against phosphorylated Syntide-2 (SEQ ID NO. 15) was raised in rabbits. The rabbit polyclonal antiphosphosyntide antibody was found to quantitatively and specifically recognize phosphoserine on both Syntide 2-Ser-PO₃ (assay on DNA BIND plates) or on biotin-Syntide 2-Ser-PO₃ (assayed on Neutravidin coated 96 well plates) when compared with the unphosphorylated peptide counterparts. A modified Chk1 HTS assay ELISA was developed using His-tagged KH289 Chk1 kinase, biotinsyntide substrate assayed on Neutravidin coated 96 well plates, and the rabbit anti-phosphosyntide antibody to detect the phosphorylated product.

[0172] This Chk1 kinase ELISA HTS allowed for the robotic screening of compound libraries. Herein, the Beckman robotics station was used. First; the Chk1 kinase was assayed in Neutravidin coated 96-well plates in 100 μ L/well of reaction mixture. The reaction mixture comprised 50 mM Tris-HCl (pH 7.5), 10 mM MgCl₂, 3 mM DTT, 400 mM NaCl, 50 μ M ATP, 10 μ M biotin-Syntide 2 peptide substrate and 10 nM Chk1 kinase (KH289). The assay was performed both with and without 20 μ M test compound. Herein, the biotin Syntide 2 substrate had the following sequence: PLARTLSVAGLPGK-biotin-K (SEQ ID NO. 12).

[0173] The assay is depicted in Figure 10. In step A, 93 μ L of reaction mixture (less both the Chk1 kinase and the biotin-syntide) is added, followed by the addition of 2 μ L of test compound (20 μ M final). The kinase reaction is initiated by the addition of 5 μ L of enzyme-substrate stock (200 nM Chk1 kinase and 200 μ M biotin-syntide). The kinase reaction is allowed to proceed for 10 min at room temperature (\approx 22 °C) as shown in Step B. Following 10 minutes of kinase reaction, both phosphorylated and unphosphorylated biotin-Syntide 2 are bound to the Neutravidin coated plate. In step C, the plates are washed with PBS/Tween-20 to terminate the kinase reaction and to remove the unbound phosphorylated or non-phosphorylated biotin-Syntide 2. In step D, the plates are incubated at room temperature for 60 minutes with rabbit anti-phosphosyntide antibody (1: 40,000 dilution; 100 μ L/well). The anti-phosphosyntide antibody binds specifically to the serine-phosphorylated biotin-Syntide 2. The unbound antibody is removed with washes of PBS/Tween-20. The plates are then incubated at room temperature for 60 minutes with goat-anti-rabbit-IgG(Fc)-HRP (horseradish peroxidase) antibody. In step E, the plates are washed with PBS/Tween to remove the unbound secondary antibody. Then, 100 μ L/well chromogenic dye ABTS (HRP substrate) is added. The color development, resulting from the HRP reaction, is allowed to take place for 18 minutes. This is followed by absorbance measurement at 405 nm in a 96-well plate reader. The Chk1 kinase activity is directly proportional to the optical density of the color formed.

[0174] All references cited herein are incorporated by reference in their entirety.

[0175] While the invention has been described in conjunction with examples thereof it is to be understood that the foregoing description is exemplary and explanatory in nature, and is intended to illustrate the invention and its preferred embodiments. Through routine experimentation, the artisan will recognize apparent modifications and variations that may be made without departing from the spirit of the invention. Thus, the invention is intended to be defined not by the above description, but by the following claims and their equivalents.

SEQUENCE LISTINGS

[0176]

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SEQ ID NO. 1 — full length human Chk1 (nucleotide sequence — 1933 base pairs)
SEQ ID NO. 2 — full length human Chk1 (peptide sequence - 476 AA)
SEQ ID NO. 3 — PCR primer (chk6w)
SEQ ID NO. 4 — PCR primer (KH289)
SEQ ID NO. 5 — PCR primer (K289)

SEQ ID NO. 5 — PCR primer (Chk11)
SEQ ID NO. 7 — PCR primer (K210)
SEQ ID NO. 8 — PCR primer (KH210)
SEQ ID NO. 9 — PCR primer (KH248)
SEQ ID NO. 10 — PCR primer (KH248)
SEQ ID NO. 11 — synthetic substrate peptide, Syntide-2
SEQ ID NO. 12 — synthetic substrate peptide, Syntide-3
SEQ ID NO. 13 — synthetic substrate peptide, Syntide-4
SEQ ID NO. 14 — oligonucleotide primer
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	EP 1 096 014 A2
	SEQ ID NO. 15 — serine phosphorylated Syntide-2
5	SEQ ID NO. 16 — serine phosphoxylated biotin Syntide-2 SEQ ID NO. 17 —peptide sequence for Cdc25 protein phosphatase SEQ ID NO. 18 —peptide sequence for mouse (mm) Chk1 kinase domain SEQ ID NO. 19 —peptide sequence for Xenopus (x1) Chk1 kinase domain SEQ ID NO. 20 —peptide sequence for fruit fly (dm) Chk1 kinase domain SEQ ID NO. 21 —peptide sequence for C. elegans (ce) Chk1 kinase domain SEQ ID NO. 22 —peptide sequence for S. cerevisiae (sc) Chk1 kinase domain
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<110> Agouron Pharmaceuticals, Inc.

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10	-	er Tyr 90	Asp	Leu	Val	Asp 295	Ser	Ala	Ala	Ala	Leu 300	Glu	lle	Cys	Ser
	Pro 305								•						
15	<210><211><211><212><213>	299	egan:	s				**							
20	<400> Met S	21 er Ala	Ala	Ser 5	Thr	Thr	Ser	Thr	Pro 10		Ala	Ala	Ala	Val 15	
	Pro G	ln Gln	P=0 20	Glu	Ser	Leu	Tyr	Arg 25	Val	Val	Gln	Thr	Leu 30	Gly.	Glu
25	Gly A	la Phe 35	_	Glu	Val		Leu 40	Ile	Val	Asn	Thr	Lys 45	Asn	Pro	Glu
		la Ala 50		Met		Lys 055		Asn	Ile	Ala	Asn 60	Lys	Ser	Lys	Asp
30		le Asp		Ile	Arg 70	_	Glu	Tyr		Leu 75	Gln	Lys	Arg	Val	Ser 80
		al Gly	:	∵ 85		·:			.90		•	•		95	
35		ln Phe	100					105	-				110	•	
:		sp Lys 115			,		120	•				125		•	
40	1	yr Phe 30				135					140				
•	145	al Val			150					155		•	•	•	160
45		is Val		165		•			170					175	
		ys Gly	180					185					190		
50	_	la Ala 195				•	200					205			
		sp Val 10	Trp	Ser	Ser	Gly 215	Ile	Val	Leu	Ile	Ala 220	Met	Leu	Thr	Gly

		Gl u 225	Leu	Pro	Trp	Asp	Arg 230		Ser	Asp	Ala	Ser 235	Gln	Seï	Týr	Met	ĞÎy 240
5		Trp	Ile	Ser	Asn	Thr 245	Ser	Leu	Asp	Glu	Arg 250		Trp	Lys	Lys	Ile 255	Asp
		Val	Arg	Ala	Leu 260	Cys	Met	Leu	Arg	Lys 265		Val	Thr	Asp	Lys 270		Asp
10		Lys	Arg	Ala 275	Thr	Ile	Glu	Gln	11e 280		Ala	Asp	Pro	Trp 285		Gln	His
		Λsn	290		Gln	Val	Glu	Thr 295		Asn	Gly	Arg					
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	·	Leu	Gly	Asp	Thr 20	Val	Gly	Gln	Gly	Ala 25	Phe	Ala	Cys	Val	Lys 30	Asn	Ala
25	:	His	Leu	Gln 35	Met	Asp	Pro	Ser	Ile 40	Ile	Leu	Ala	Val	Lys 45	Phe	Ile	His
		Val	Pro 50	Thr	Cys	Lys	Lys	Met 55	Gly	Leu	Ser	Asp	Lys 60	Asp	Ile	Thr	Lys
30		65		٠.			70		•			75					Arg - 80
·		٠			Cys	85				•	90					95	
35					Gly 100					105					110		
				115	Asp				120					125			
40			130		Leu			135					140				
•	:	145			Ile Ala		150					155					160
45		•				165				•	170					175	
		•			Gln 180 Glu					185				٠	190		
50				195	Leu				200				•	205		·	
		C1 y	116	Dou	200		<i>,</i>		200	4114 .	., L Y	O T 11	THE	L10	пр	φŧū	rea

			210					215	• •				⁻ 220		• =	••	
5		Pro 225	Ser	Leu	Glu	Asn	Glu 230	Asp	Phe	Vaļ	Phe	Phe 235	Ile	Glu	Asn	Asp	Gly 240
		Asn	Leu	Asn	Trp	Gly 245	Pro	Trp	Ser	Lys	Ile 250	Glu	Phe	Thr	His	Leu 255	Àsn
10		Leu	Leu	Arg	Lys 260	Ile	Leu	Gln	Pro	Asp 265	Pro	Asn	Ļys	Arg	Val 270	Thr	Leu.
		Lys	Ala	Leu 275	Lys	Leu	His	Pro	Trp 280	Val	Leu	Arg	Arg	Ala 285	Ser	Phe	Ser
15		Gly	Asp 290	Asp	Gly	Leu	Cys	Asn 295	Asp	Pro	Glu	Leu	Leu 300	Ala	Lys	Lys	Leu
		Phe 305	Ser						•				٠.	,			
20	·	<21 <21	0> 2: 1> 2: 2> P: 3> S	95 R T	nbe							•		· .		* 2	
25			0> 2: Ala		Lys	Leu 5	Asp	Asn	Phe	Pro	Tyr 10	His	Ile	Gly	Arg	Glu 15	Ile
:		Gly	Thr	Gly	Ala 20	Phe	Ala	Ser	Val	Arg 25	Leu	Cys	Tyr	Asp	Asp 30	Asn	Ala
30 ·		Lys	Ile	Tyr 35	Ala	Val	Lys	Phe	Val .40		Lys	Lys		Ala 45	Thir	Ser	Cys
		Met	Asn 50	Ala	Gly	Val	Trp	Ala 55	Arg	Arg	Met	Ala	Ser 60	Glu	Ile	Gln.	Leu
35		His 65	Lys	Leu	Cys	Asn	Gly 70	His	Lys ·	Asn	Ile	11e 75	His	Phe	Ţyr	Asn	Thr 80
		Ala	Glu	Asn	Pro	Gln 85	Trp	Arg	Trp	Val	Val 90	Leu	Glu	Phe	Ala	.Gln 95	Gly
40	•				100					105					Asp 110		
				115					120					125		٠.	Met ·
45			130					135				. •	140		Asn		
		145	_	-			150					.155			Phe		160
50		Leu	Phe	Ser	Tyr	Lys 165	Gly	Lys	Ser	Arg	Leu 170	Leu	Asn	Ser	Pro	Val 175	Gly
		Ser	Pro	Pro	Tyr 180	Ala	Ala	Pro	Glu	Ile 185		Gln	Gln	Tyr	Asp 190	Gly	Ser

```
Lys Val Asp Val Trp Ser Cys Gly Ile Ile Leu Phe Ala Leu Leu Leu
                    195
                                                             205
                                         200
           Gly Asn Thr Pro Trp Asp Glu Ala Ile Ser Asn Thr Gly Asp Tyr Leu
                                    215
           Leu Tyr Lys Lys Gln Cys Glu Arg Pro Ser Tyr His Pro Trp Asn Leu
                                230
                                                                          240
           225
                                                     235
10
           Leu Ser Pro Gly Ala Tyr Ser Ile Ile Thr Gly Met Leu Arg Ser Asp
                                                 250
           Pro Phe Lys Arg Tyr Ser Val Lys His Val Val Gln His Pro Trp Leu
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                                             265
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40
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45
```

Claims

- A composition comprising an isolated, purified polynucleotide which encodes the active form of the human Chk1 kinase or a functional, active human Chk1 kinase analog thereof.
- 2. The composition according to claim 1, wherein the nucleotide sequence of said polynucleotide comprises bases 35 to 830 of SEQ ID NO. 1 or a functional, active mutant or variant thereof.
 - 3. A polypeptide in a crystallized form comprising the catalytically active form of the human Chk1 kinase and the inhibitor binding site thereof.

- The polypeptide according to claim 3 wherein the crystal is solved to a resolution of at least 2.5 ().
- The polypeptide according to claim 3 wherein the crystal is solved to a resolution of at least 2.0 ().
- The polypeptide according to claim 3 wherein the crystal is solved to a resolution of about 1.7 ().

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- 7. The polypeptide according to claim 3 wherein the amino acid sequence of said polypeptide comprises amino acids 16 to 265 of SEQ ID NO. 2 or an active mutant or variant thereof.
- The polypeptide according to claim 3 wherein the amino acid sequence of said polypeptide comprises amino acids 16 to 289 of SEQ ID NO. 2 or an active mutant or variant thereof.
 - 9. The polypeptide according to claim 3 wherein the amino acid sequence of said polypeptide comprises amino acids 16 to 291 of SEQ ID NO. 2 or an active mutant or variant thereof.
 - 10. The polypeptide according to claim 3 wherein the amino acid sequence of said polypeptide comprises amino acids 1 to 265 of SEQ ID NO. 2 or an active mutant or variant thereof.
 - 11. The polypeptide according to claim 3 wherein the amino acid sequence of said polypeptide comprises amino acids 1 to 289 of SEQ ID NO. 2 or an active mutant or variant thereof.
 - 12. The polypeptide according to claim 3 wherein the amino acid sequence of said polypeptide comprises amino acids 1 to 291 of SEQ ID/NO. 2 or an active mutant or variant thereof.

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- 13. The polypeptide according to claim 3 wherein said polypeptide further comprises a six histidine tag on the: C-terminal thereof. Could be thanked the control of the transfer of the control of the
 - 14. An isolated, soluble, catalytically active polypeptide comprising the active form of the human Chk1 kinase or a functional, active human Chk1 kinase analog thereof.
 - 15. The polypeptide according to claim 14 comprising the full length human Chk1 protein having the C-terminal portion thereof deleted so as yield the human Chk1 kinase domain in its active configuration.
 - 16. The polypeptide according to claim 14 wherein said polypeptide comprises amino acids 16 to 265 of the sequence as set forth in SEQ ID NO. 2 or a conservatively substituted variant thereof.
 - 17. The polypeptide according to claim 14 wherein said polypeptide comprises amino acids 16 to 289 of the sequence as set forth in SEQ ID NO. 2 or a conservatively substituted variant thereof.

- 18. The polypeptide according to claim 14 wherein said polypeptide comprises amino acids 16 to 291 of the sequence as set forth in SEQ ID NO. 2 or a conservatively substituted variant thereof.
 - 19. The polypeptide according to claim 14 wherein said polypeptide comprises amino acids 1 to 265 of the sequence as set forth in SEQ ID NO.2 or a conservatively substituted variant thereof.
 - 20. The polypeptide according to claim 14 wherein said polypeptide comprises amino acids 1 to 289 of the sequence as set forth in SEQ ID NO. 2 or a conservatively substituted variant thereof.
- 21. The polypeptide according to claim 14 wherein said polypeptide comprises amino acids 1 to 291 of the sequence as set forth in SEQ ID NO. 2 or a conservatively substituted variant thereof.
 - 22. The polypeptide according to claim 14 wherein said polypeptide comprises amino acids 5 to 265 of the sequence as set forth in SEQ ID NO. 2 or a conservatively substituted variant thereof.
- 23. The polypeptide according to claim 14 wherein said polypeptide comprises amino acids 5 to 289 of the sequence as set forth in SEQ ID NO. 2 or a conservatively substituted variant thereof.
 - 24. The polypeptide according to claim 14 wherein said polypeptide comprises amino acids 5 to 291 of the sequence

as set forth in SEQ ID NO. 2 or a conservatively substituted variant thereof.

- 25. An expression vector for producing active human Chk1 kinase in a host cell, which vector comprises: a polynucleotide encoding active form of the human Chk1 kinase or an active human Chk1 kinase analog thereof; transcriptional and translational regulatory sequences functional in said host cell operably linked to said human Chk1 kinase-encoding polynucleotide; and a selectable marker.
- 26. The vector according to claim 25 wherein said polynucleotide encodes the active human Chk1 kinase, said active kinase comprising bases 35 to 830 of SEQ ID NO. 1.
- 27. The vector according to claim 25 wherein said vector is selected from the group consisting of pET28a, pAcSG2, and pFastBac.
- 28. The vector according to claim 25 wherein said vector is pFastBac-Nde.
- 29. The vector according to claim 25 wherein said selectable marker is selected from the group consisting of beta galactosidase, green fluorescent protein, and luciferase.
- 30. A host cell stably transformed and transfected with a polynucleotide encoding active form of the human Chk1 kinase or an active human Chk1 kinase analog thereof in a manner allowing the expression in said host cell of the human Chk1 kinase.
 - 31. The host cell according to claim 30, wherein said polynucleotide encodes the active hChk1 kinase, said active kinase comprising bases 35 to 830 of SEQ ID NO. 1.
 - 32. The host cell according to claim 30 wherein said host is E. coli.
 - 33. The host cell according to claim 30 wherein said host is a recombinant baculovirus.
- 30 34. The host cell according to claim 30 wherein said host is an insect cell.
 - 35. The host cell according to claim 34 wherein said insect cell is Sf9.
 - 36. The host cell according to claim 30 wherein said host cell is transformed and transfected with said polynucleotide via an expression vector comprising said polynucleotide; a transcriptional and translational regulatory sequences functional in said host cell operably linked to said hChk1 kinase-encoding polynucleotide; and a selectable marker.
 - 37. The host cell according to claim 36 wherein said expression vector is selected from the group consisting of pET28a, pAcSG2, and pFastBac.
 - 38. The host cell according to claim 36 wherein said expression vector is pFastBac-Nde.
 - 39. The host cell according to claim 36 wherein said selectable marker is selected from the group consisting of beta galactosidase, green fluorescent protein, and luciferase.
 - 40. A method for assaying a candidate compound for its ability to interact with the human Chk1 comprising:
 - (a) expressing an isolated DNA sequence or variants thereof encoding the kinase domain of said human Chk1 in a host capable of producing said kinase in the catalytically active configuration, said kinase in a form which may be assayed for interaction of said kinase with said candidate compound;
 - (b) exposing said kinase to said candidate compound; and
 - (c) evaluating the interaction of said kinase with said candidate compound.
 - 41. A method of identifying a Chk1 kinase inhibitor by determining the binding interactions between an organic compound and the binding site of the Chk1 kinase in the active conformation, said binding sites being defined by the crystal coordinates of provided in Figure 11, said method comprising:
 - (a) generating the binding cavity defined by the binding site on a computer screen;

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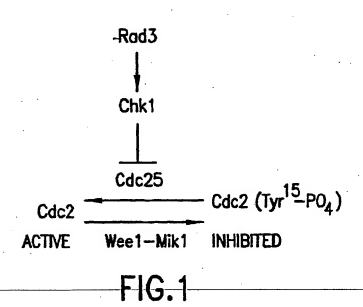
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40

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	(h) generating compo	unds with their spatial structure;	and	
	(c) testing to see whe	ther the compounds bind to at t ing site can be identified as Chk	he Chk1 binding site; wherein	those compounds that do
5				
		•		
10				
		•		
15		,		
20				
	·	; ;	1. N. A.	(1.5) (25) (3.5)
<i>25</i> '				
20		·.		* 2.# %
		*		₹\ *
<i>30</i>				Ξ.
35				
ω		Single Si		
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40				
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43			·	



55 55 55 27 167 154 166 <u>ज</u> IEPDICAPECDACRE FHOLMACYVYLIG-I GITHRDIKPENILLD ERDNIKISDFGLATV EPOVGMPECDACKE FOOL JACVEYLHS-1 GITHRD JKPENLLLD EROCLKISDFCLATV OTCCCCGMANDLKK ---- HPDAANSVR KEVC10MLO--DK ce nsaastistpaaaav apoopeslyrvvotl gegafgevilivntk npevaaakkinian ---- kskofidnir keyllokrysavchd NAVKFYCHRRECNIQ YLFLEYCSCCELFDR IEPDIGMPEPDAQRF FHQLMACWYLHS-I GITHRDIKPENLLLD ERDNLKISDFGLATV AVDCPENIK KEICINKMIN--HE AIDCPONIK KEICINAMES--HE --- m acklonepyhigrei gigafasyricyd--dnakiyavkfvnnkh atschaugcwarrwa seiglhkicn--ghk EPONOMPONEAGRY FIGUL SCLIMING-R CLANROLKPENLLLO ENDIVIKISOFCIMATIN EPOCONCENT LLT GTHVLKISOFCHATL EPDYGYDSDYAGFY FOOLVSAINMINEC GVAHPDIKPENILLD KNONLKLADFGLASO EPDVGIDEDVAQFY FAQLMEGISFMAS—K GVAHROLKPENIULD YNGNLKISDFGFASI AADCPENIK KEICINRALS--H දි I TEDAVANK I VOJAKR GEGAYCEVOLAVN-R KTEEAVAVKIVDMTR ----GEGAYGEVOLAVN-R VTEEAVAVKIVDMKR ---岁 GEGAYCEVOLAVN-R GEGAYGEVKLL IN-R NVVKFYGHRREGHIO YLFLEYCSGGELFDR NIVRFYCHRREGNIQ YLFLEYCRGGELFDR HILRFECKRSGGSVE YIFLEYAAGGELFDR NVLRL I DCNVSKEYM WI I LEMADGGDLF DK NITHEYNTAENPOWR WAVLEFACCOLFDK YLFLEYADGGELFDK REFVECINTLAQTL MAYPFYEDWDL YOTL WAVPF VEDMOL VOTL AVPF VEDWOL VQTL 늉 NV I RAJ CARRADPOFY dim MAATL TEACTGPAAT E Sp SC S ş × 튱 ဗ္ဗ

GTLPYVAPELLKR-K EFHAEPVDVMSCGIV LTAM AGELPMOQPS OSCOEYSDMKEK--K TYLNPMKKIDSAPLA 240 GTLPYVAPEVLO--K AYOPOPADLWSGGVI LYTMLAGELPMOOPS TNCTEFTNWRONDHW OLOTPWSKLDTLAIS 254 EVCOEYCONKEK -- N HYLTPWKK [SATPLA DASQSYMCWISN-TS LDERPWKK I DVRALC GSPPYMAPEVLYSEE GYYADRIDINSIGIL LFVLLIGOTPHELPS LENEDFVFFIENDON LINIGPHSKIEFTHLN GTLPYVAPELLKR-K EFHAEPVOVNSOCIV LTAM, AGELPHDOPS DSCOEYSDWKEK--K TYLNPWKKIDSAPLA SNIGDYLL YKKOCER PSYHPWALL SPGAYS GTLPYVAPELIKS-R AFHADPYDVMSCGIV LIAMLAGELPMDQPN GTIPYAAPELCAG-K KYRCPPYDVWSSGIV LIAM TGELPWDRAS GSPPYAAPE 110--- QYDCSKYDVMSCC11 LFALLLGNTPMDEA1 FRENCHIOL FRIN-CKER LSKAC FRRF DCT LRVSNOOR FRYN-NRERLLINGIC FRCK-CKERLLDKRC YRNK-GEERLLOLSC E ş ≂ 통 ೮

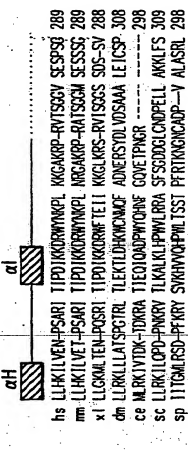


FIG 2R

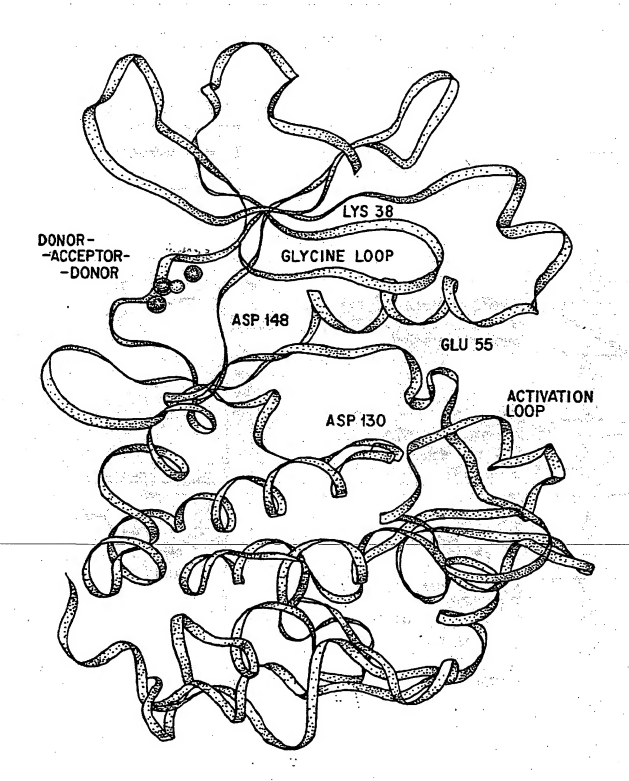


FIG. 3

His-tagged CHK1 Kinase domain 1-289 Purification

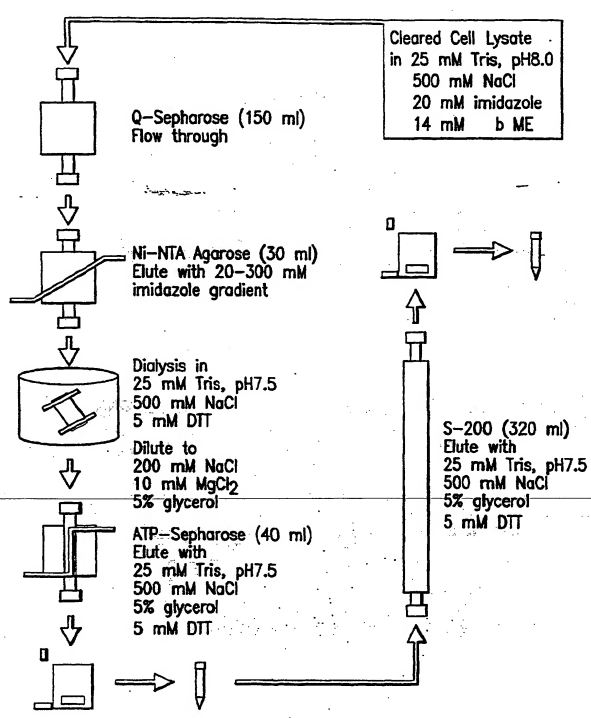


FIG.4

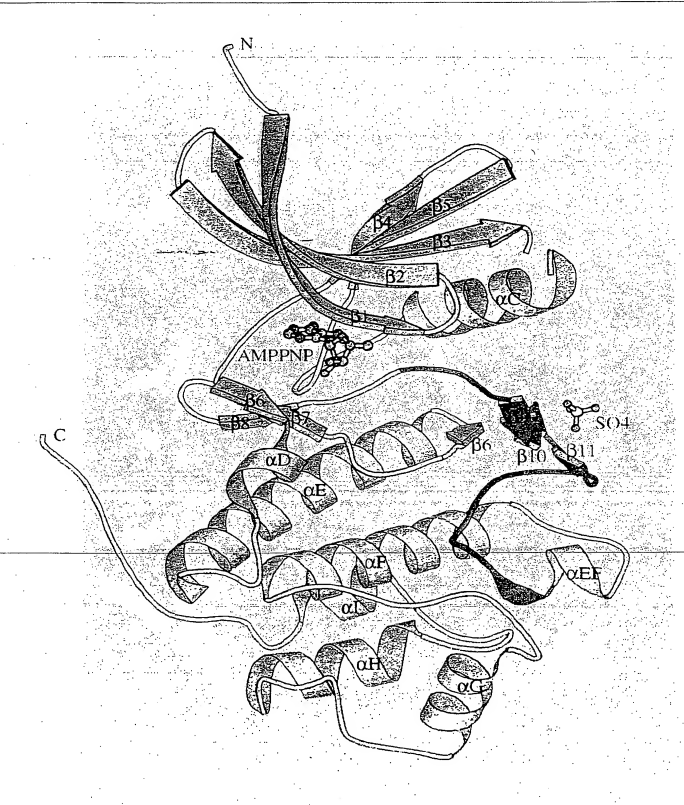
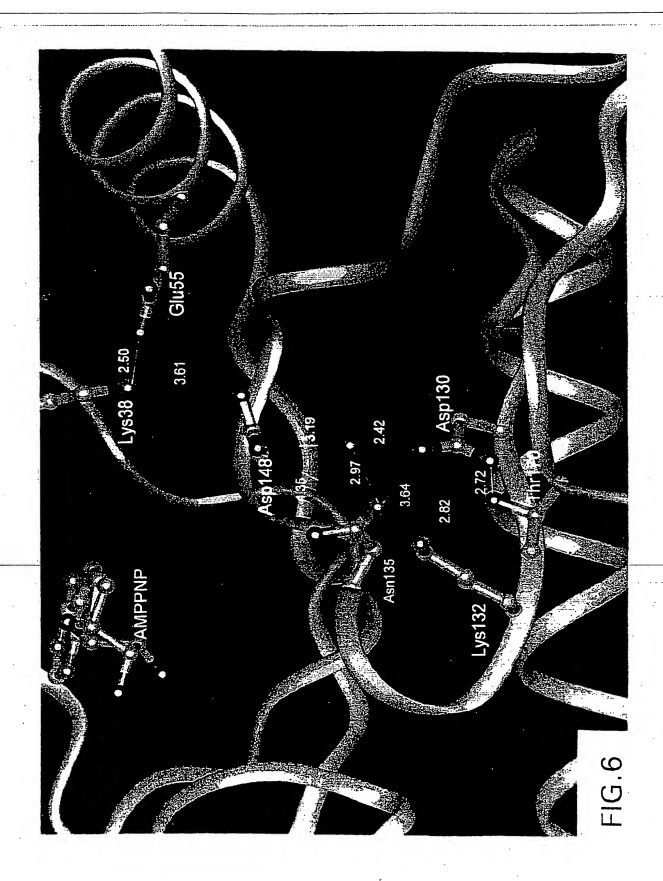
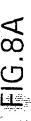
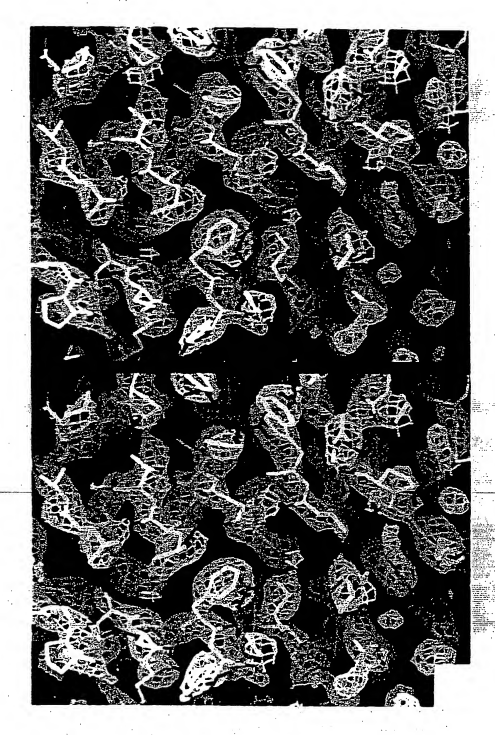


FIG.5









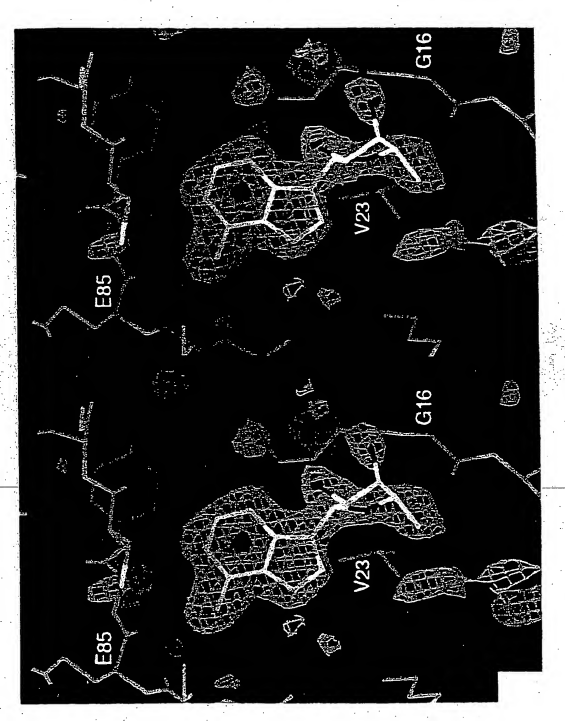
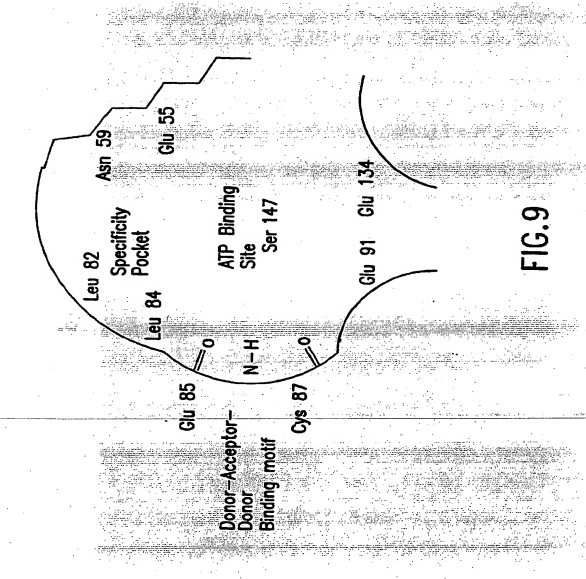
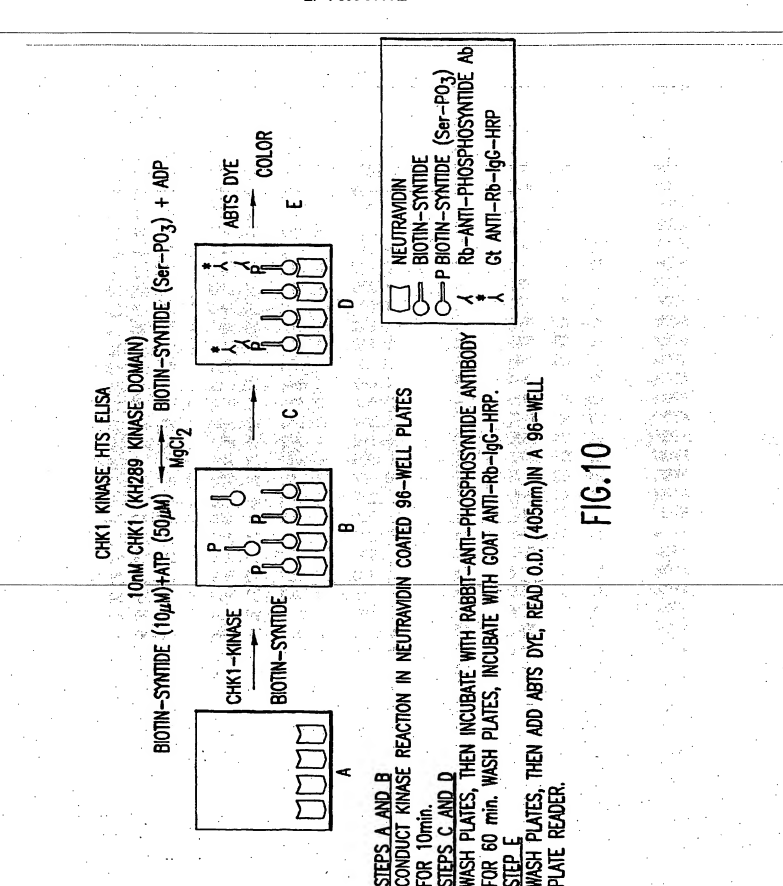


FIG. 8B





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ATOM	3	0	ALA	2	0.890		-14.560	1.00 54.24
ATOM	4	N	ALA	2	-0.778		-11.709	1.00 57.23
ATOM	5	CA	ALA	2	-0.258		-12.949	1.00 55.69
ATOM	6	N	VAL	3	2.056		-12.740	1.00 51.32
ATOM:	7	· CA	VAL	3	3.284		-13.149	1.00 47.11
ATOM	8	CB	VAL	3	4.508		-12.363	1.00 46.10
MOTA	9	CG1	VAL	3	5.794		-12.973	1.00 41.34
ATOM	10	CG2	VAL	3	4.524		-12.367	1.00 48.87
ATOM	11	C	VAL	3	3.143		-12.922	1.00 44.85
ATOM	12	0	VAL	3	2.969		-11.795	1.00 45.58
MOTA	13	N	PRO	4	3.231		-14.003	1.00 41.89
ATOM	14	CD	PRO	4	3.546		-15.363	1.00 37.40
ATOM	15	CA	PRO	4	3.112		-13.991	1.00 41.11
ATOM	16	CB	PRO	-4	3.743		-15.323	1.00 34.82
ATOM	17	CG	PRO	4	3.281	4.388		1.00 31.95
ATOM	18	C	PRO	4	3.667		-12.815	1.00 42.75
ATOM	19	0	PRO	4	2.936		-11.875	1.00 47.35
ATOM	20	N	PHE	5	4.954	2.540	-12.869	1.00 40.95
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MOTA	24	CD1	PHE	5	7.335		-14.862	1.00 27.57
ATOM	25		PHE	5	5.702	-0.154	-14.589	1.00 32.16
ATOM	26		PHE	5	7.237	1.465	-16.248	1.00 27.10
ATOM	27		PHE	5	5.593	-0.276	-15.979	1.00 30.91
ATOM	28	CZ	PHE	5	6.363	0.535	-16.809	1.00 28.05
ATOM	29	-C	PHE	5	6.156	2.348	-10.589	1.00 39.99
ATOM	30	0 -	PHE	5 6	7.191	1.908	-10.088	1.00 38.49
ATOM	31	N	VAL		5.486	3.360	-10.048	1.00 40.37
ATOM	32		VAL	6	5.994	4.011	-8.842	1.00 40.35
ATOM	33	CB	VAL	6	5.424	5.437	-8.690	1.00 42.42
ATOM	34		VAL	6	6.135	6.169	-7.563	1.00 45.17
ATOM	35		VAL	6	5.593	6.194	-9.980	1.00 43.26
ATOM	36		VAL	6	5.676	3.219	-7.573	1.00 39.11
ATOM	37	0	VAL	6	6.229	3.492	-6.507	1.00 38.75
ATOM	38	N	GLU	7	4.796	2.232	-7.693	1.00 36.63
ATOM	39	CA	GLU	7	4.408	1.411	-6.550	1.00 34.52
ATOM	40	CB	GLU	7	2.931		-6.659	1.00 41.81
ATOM	41	CG	GLU	7	1.981	2.219	-6.618	1.00 51.32
ATOM	42	CD	GLU	7	1.963	2.906	-5.267	1.00 60.70

FIG.11A-1

MOTA	43	0E1	GLU	7 .	3.021	3.416	-4.840	1.00 63.96
ATOM	44	OE2	GLU	7	0.888	2.935	-4.631	1.00 70.24
MOTA	45	C	GLU	7	5.246	0.141	-6.443	1.00 31.93
MOTA	46	0	GLU	7	5.036	-0.675 ·	-5.544	1.00 32.27
MOTA	47	N .	ASP	8	6.193	-0.024	-7.360	1.00 31.04
MOTA	48	CA	ASP	8	7.052	-1.204	-7.367	1.00 31.42
MOTA	49	CB .	ASP	8	7.404	-1.597	-8.805	1.00 35.69
MOTA	50	CG .	ASP	8 4 4	6.202	-2.095		1.00 44.50
MOTA	51	OD1	ASP	8	5.534	-3.039	-9.115	1.00 46.19
MOTA	52	OD2	ASP	8	5.929	-1.544	-10.673	1.00 49.49
ATOM	53	C	ASP	8 ****	8.338	-1.002	-6.576	1.00 31.44
MOTA	54	0	ASP	8	9.039	-0.003		1.00 32.04
ATOM	一 55	N	TRP	9	8.644	-1.972	-5.720	1.00 30.97
ATOM	56	CA	TRP	9	9.837	-1.939	-4.883	1.00 31.19
MOTA	57	CB	TRP	9	9.435	-1.779		1.00 32.92
MOTA	58	CG	TRP	9	9.527	-0.371		1.00 37.61
ATOM	59	CD2	TRP	9	8.464	0.583	-2.781	1.00 34.10
MOTA	60	CE2	TRP	9	9.014	1.770	-2.242	1.00 35.49
MOTA	61	CE3	TRP	9	7.100	0.554	-3.100	1.00 28.78
MOTA	62	CD1	1.5	9	10.648	0.258	-2.404	1.00 37.08
MOTA	63			9	10.347	1.543	-2.020	1.00 32.55
ATOM	64	CZ2		9	8.247		-2.015	1.00 34.87
ATOM	65			9	6.337	1.693	-2.876	1.00 36.67
MOTA		CH2			6.914	2.860	-2.338	1.00 40.32
ATOM	67		TRP	9 💝 😕	10.666	-3.213	-5.044	1.00 31.47
MOTA	68		TRP	9	10.127	-4.320	-5.057	1.00 32.63
MOTA	69		ASP	10	11.977	-3.046	-5.173	1.00 30.81
MOTA			ASP_	10	12.893		5.304_	_1.00_30.35_
MOTA	71		ASP	10	13.994		-6.316	1.00 31.71
MOTA	72		ASP	10	13.474	-3.745	-7.736	1.00 35.37
MOTA		OD1		10	14.061	-2.971		1.00 41.94
MOTA		OD2		10	12.495	-4.445	-8.069	1.00 34.42
MOTA		• • • • • • • • • • • • • • • • • • • •	ASP	10	13.539			1.00 30.57
MOTA	76		ASP	10	14.029	-3.521		1.00 27.52
MOTA	77		LEU	11	13.535		-3.522	1.00 32.48
MOTA	78		LEU	11	14.148	-6.078	-2.249	1.00 36.27
MOTA	79		LEU	11	13.432	-7.290	-1.645	1.00 37.99
MOTA	80		LEU	11	11.990	-7.058	-1.182	1.00 39.05
ATOM	81	CD1		11	11.125	-6.630	-2.357	· - -
MOTA	82	CD2		11	11.442	-8.335	-0.563	1.00 43.43
MOTA	83		LEU	11	15.609	-6.405	-2.537	
MOTA	84	0.	LEU	11	15.934	-7.508	-2.975	1.00 38.62

FIG.11A-2

ATOM	85	-N	VAL	12	16.480	-5.432	-2.287	1.00 41.77
MOTA	86	CA	VAL	12	17.909	-5.563	-2.557	1.00 44.63
ATOM	87	CB	VAL	12	18.555	-4.169	-2.720	1.00 45.32
ATOM	88	CG1	VAL	12	20.017	-4.310	-3.124	1.00 51.19
ATOM	89	CG2	VAL	12	17.788	-3.365	-3.757	1.00 42.65
ATOM	90	C	VAL	12	18.739	-6.353	-1.549	1.00 46.95
ATOM .	91	0	VAL	12	19.663	-7.068	-1.937	1.00 47.15
ATOM	92	N	GLN	13	18.431	-6.223	-0.262	1.00 47.33
MOTA	93	CA	GLN	13 .	19.195	-6.940	0.752	1.00 47.97
ATOM	94	CB	GLN	13	20.558	-6.275	0.948	1.00 49.67
MOTA	9 5	CG	GLN	13	20.482	-4.801	1.303	1.00 53.58
ATOM	96	CD	GLN	13	21.833	-4.223	1.675	1.00 55.37
ATOM	97	OE1	GLN	13	22.410	-4.578	2.703	1.00,56,20
MOTA	98	NE2	GLN	13	22.347	-3.329	0.836	1.00 58.05
MOTA	99	C	GLN	13	18.505	-7.055	2.104	1.00 48.35
MOTA	100	0	GLN	13	17.636	-6.255	2.452	1.00 47.74
MOTA	101	N	THR	14	18.916	-8.063	2.866	1.00 48.79
ATOM	102	CA	THR	14	18.365	-8.310	4.192	1.00 49.84
ATOM	103		THR	14	18.497	-9.795	4.575	1.00 51.16
ATOM	104	0G1		14	18.202	-9.961	5.968	1.00 53.80
ATOM	105	CG2		14	19.907	-10.293	4.293	1.00 55.63
ATOM	106		THR	14	19.106	-7.478	5.229	1.00 49.90
MOTA	107		THR	14	20.334	-7.512	5.293	1.00 51.70
ATOM	108		LEU	15	18.363	-6.726	6.034	1.00 48.90
ATOM	109		LEU	15	18.977	-5.903	7.067	1.00 49.63
ATOM	110	•	LEU	15	18.139	-4.650	7.344	1.00 44.87
ATOM	111		LEU	15	17.959	-3.650	6.203	1.00 39.00
ATOM	112	CD1		15	19.307	-3.313	5.581	1.00 32.83
ATOM	-113 -	CD2		15	17.039	-4.247	5.172	1.00 41.59
ATOM	114		LEU	15	19.120	-6.706	8.349	1.00 51.59
ATOM	115		LEU	15	20.050	-6.493	9.126	1.00 51.40
ATOM	116	-	GLY		18.191		8.562	1.00 53.10
ATOM	117		GLY.		18.227	-		
ATOM	118	C		16	17.043		9.824	
MOTA	119		GLY	16	15.909		9.550	1.00 58.89
ATOM	120		GLU	17		-10.651	10.191	1.00 60.68
ATOM	121	CA		17	16.257		10.301	1.00 63.27
ATOM	122		GLU	17		-12.961	9.644	1.00 67.17
ATOM			GLU	17		-12.845	8.156	1.00 69.72
MOTA			GLU	17		-14.155		1.00 74.27
ATOM		.0E1		17		-14.672		1.00 77.30
ATOM	126	0E2 (GLU	17	16.727	-14.670	6.653	1.00 75.48

FIG.11A-3

· · · · · · · · · · · · · · · · · · ·							
ATOM	127	С	GLU	17	15.914 -11.911	11.762	1.00 64.57
MOTA	128	0	GLU	17	16.591 -12.682	12.441	1.00 63.90
MOTA	129	N	GLY	18	14.859 - 11.258	12.238	1.00 66.47
MOTA	130	CA	GLY	18	14.445 -11.429	13.618	1.00 66.94
MOTA	131	С	GLY	18	13.834 -12.793	13.870	1.00 67.94
ATOM	132	0	GLY		13.610 -13.565	12.936	1.00 68.20
MOTA	133	N	ALA	19	13.565 -13.093	15.137	1.00 68.69
MOTA	134	CA	ALA	19	12.973 -14.370	15.512	1.00 67.96
MOTA	135	CB	ALA	19	13.110 -14.586	17.015	1.00 67.41
ATOM	136	C	ALA	19	11.504 -14.412	15.107	1.00 67.21
ATOM	137	0.	ALA	19	10.812 -15.403	15.346	
ATOM	138	N	TYR	20	11.035 -13.330	14.493	1.00 66.16
MOTA	139		TYR	20	9.648 -13.236	14.052	1,00 65.86
MOTA	140	CB	TYR	20	8.813 -12.492	15.101	1.00 66.27
ATOM	141	CG	TYR	20	9.495 -11.278	15.697	1.00 68.01
ATOM	142		TYR	20	9.896 -10.210	14.894	1.00 72.07
ATOM	143		TYR		10.528 -9.093	15.442	1.00 72.81
ATOM	144		TYR	20	9.743 -11.201	17.068	1.00 64.75
ATOM	145		TYR	20	10.373 -10.090	17.625	1.00 66.10
ATOM	146		TYR	20	10.762 -9.041	16.806	1.00 71.06
MOTA	147	OH	TYR	20	11.385 -7.942		1.00 74.54
MOTA	148	С	TYR	20	9.522 -12.549		
ATOM	149	0	TYR		8.770 -11.586		1.00 63.94
MOTA	150	N	GLY		10.261 -13.058		1.00 63.95
MOTA	151	CA	GLY	21	10.222 -12.488		1.00 62.81
ATOM	152	C	GLY	21	11.583 -12.006	9.915	1.00 61.39
ATOM	153	0	GLY	21	12.616 -12.527		
ATOM	154	<u>N</u>	GLU	22	11.587 -11.008	9.038	1.00 58.87
ATOM	155	CA	GLU	22	12.831 -10.455		1.00 55.14
MOTA	156		GLU	22	13.362 -11.322	7.373	1.00 58.20
	157			22	12.435 -11.395	6.170	1.00 64.23
MOTA	158	CD	GLU	22	13.021 -12.200		1.00 70.83
	159		GLU	22	12.352 -12.322		1.00 70.63
MOTA		0E2		22	14.152 -12.711	5.174	1.00 75.29
MOTA	161	C	GLU	22	12.620 -9.032	8.018	1.00 49.83
MOTA	162	0		22	11.492 -8.610	7.773	1.00 48.81
MOTA	163	N	VAL		13.716 -8.296	7.875	1.00 47.61
MOTA		CA	VAL	23	13.656 -6.925	7.393	
MOTA	165	CB	VAL.		14.211 -5.937	8.441	1.00 43.86
ATOM	166	CG1		23	14.076 -4.512	7.935	1.00 42.93
ATOM	167	CG2		23	13.469 -6.107	9.756	1.00 40.58
ATOM	168	C	VAL	23	14.479 -6.819	6.117	1.00 40.96

FIG.11A-4

					*			
MOTA	169	0	VAL	23	15.651	-7.190	6.091	1.00 38.29
ATOM	170	N	GLN	24	13.853	-6.322		1.00 38.29
MOTA	171	CA	GLN	24	14.518	-6.172		1.00 40.30
ATOM	172	CB	GLN	24	13.749	-6.938		1.00 40.50
ATOM	173	CG	GLN	24	13.812	-8.450		1.00 47.92
MOTA	174	CD	GLN	24	15.194	-8.999		1.00 47.92
ATOM	175	0E1	GLN	24	15.789	-8.701	-	1.00 59.08
ATOM .	176	NE2	GLN	24	15.712	-9.810		1.00 59.08
ATOM	177	C	GLN	24	14.634	-4.711		1.00 39.81
ATOM	178	0	GLN	24	13.757	-3.896	_	1.00 39.30
MOTA	179	N	LEU	25	15.733	-4.387		
ATOM	180	CA	LEU	25	15.952	-3.036		
MOTA	181		LEU	25-	17.449	-2.765		1.00 36.35
ATOM	182		LEU	25	17.903	-1.405		1.00 33.98
ATOM	183	CD1	LEU	25	17.676	-1.327		
ATOM	184	CD2		25	17.159	-0.293		1.00 37.28
ATOM	185		LEU	25	15.245	-2.983	0.843	1.00 37.37
ATOM	186		LEU	25	15.589	-3.731		1.00 34.98 1.00 34.14
ATOM	187		ALA.	26	14.249	-2.111	0.733	
ATOM	188	CA	ALA	26	13.485	-1.976	-0.501	1.00 33.58
ATOM	189		ALA	26	11.996	-2.034	-0.195	1.00 32.06
ATOM	190		ALA	26	13.816	-0.682		1.00 31.61 1.00 30.85
ATOM	191		ALA	26	13.860	0.386	-0.624	1.00 30.61
ATOM	192	an I	VAL	27	14.047	-0.788	-2.535	1.00 30.81
ATOM	193		VAL	27	14.366	0.373	-3.353	1.00 29.88
ATOM	194		VAL	27	15.735	0.207	-4.046	1.00 28.11
ATOM	195	CG1 \	VAL	27	16.053	1.442	-4.877	1.00 23.89
ATOM	196	CG2 V	VAL	27	16.818	-0.016	-2.997	1.00 23.69
ATOM	197	C1	VAL	27	13.277	0.540	-4.404	1.00 25.70
ATOM	198	0. \	VAL	27	12.933	-0.409	-5.112	1.00 26.38
ATOM	199	N A	ASN	28	12.724	1.745	-4.493	1.00 23.90
ATOM	200	CA A	ASN	28	11.657	2.014	-5.444	1.00 23.36
ATOM	201	CB A	ASN	28 -	11.047	3.391	-5.185	1.00 22.07
ATOM	202	CG A	SN	28	9.822	3.652		1.00 23.58
ATOM	203	0D1 A	ASN	28	9.925	4.068		1.00 23.59
ATOM	204	ND2 A	ASN	28	8.648	3.396	-5.462	1.00 23.39
ATOM	205	C A	ISN	28	12.169	1.926	-6.872	1.00 23.17
ATOM	206	0 4	SN	28	13.204	2.493	-7.212	1.00 23.17
ATOM	207	N A	I RG	29	11.427	1.197	-7.693	1.00 21.97
ATOM	208	CA A	I RG	29	11.771	0.981	-9.094	1.00 25.26
MOTA	209	CB A	IRG	29	10.695	0.099	-9.728	1.00 25.06
ATOM	210		IRG	29	10.782		-11.235	
						U. UTT	11. LUU	1.00 22.45

FIG.11A-5

		<u>.</u>					
ATOM	211	CD ARG	29	9.652	-0.930	-11.737	1.00 20.20
ATOM	212	NE ARG	29	9.593		-13.198	1.00 19.85
ATOM	213	CZ ARG	29	8.731	-1.680	-13.901	1.00 21.65
MOTA	214	NH1 ARG	29	7.847	-2.449	-13.281	1.00 26.57
MOTA:	215	NH2 ARG	29	8.756	-1.642	-15.227	1.00 23.50
MOTA	216	C ARG	29	11.938	2.269	-9.901	1.00 25.06
MOTA	217	O ARG	29	12.784	2.347	-10.799	1.00 25.77
ATOM	218	N VAL	30	11.136		-9.576	1.00 23.54
ATOM	219	CA VAL	30	11.178	4.548	-10.291	1.00 22.97
ATOM	220	CB VAL	30	9.753	5.109	-10.499	1.00 22.15
ATOM	221	CG1 VAL	30	9.824	6.517	-11.081	1.00 23.25
MOTA	222	CG2 VAL	30.74	8.956	4.190	-11.413	1.00 20.64
ATOM	223	C VAL	30	12.014	5.635	-9.623_	1.00 24.22
MOTA	224	0 VAL	30	12.907	6.210	-10.244	1.00 24.96
MOTA	225	N THR	31	11.724	5.915	-8.355	1.00 25.29
MOTA	226		31	12.427	6.970	-7.633	1.00 25.85
ATOM	227	CB THR	31	11.537	7.554	-6.528	1.00 29.34
ATOM	228	OG1 THR	. • •	11.357	6.574	-5.498	1.00 30.34
ATON		CG2 THR	31	10.177	7.945	-7.093	1.00 32.37
ATOM	230	C THR	31	13.742	6.557	-6.989	1.00 25.05
ATOM	231	O THR	31	14.588		-6.695	1.00 24.93
ATOM	232	N GLU	32	13.901	5.256	the state of the second state of	1.00 23.56
ATON	233	CA GLU	32	15.088	4.702		1.00 25.89
ATOM			. 32	16.360	5.169	-6.855	1.00 31.18
ATOM	235		32	16.441	4.626	-8.275	1.00 36.10
ATOM	236		JE	17.781	4.857		1.00 40.49
ATON	237		32	18.800	4.385	-8.381	1.00 47.18
MOTA	238	OE2 GLU	32	17.812		<u>-9.992</u>	_1.00_34.21_
ATON	239		32	15.125	5.060		1.00 28.39
ATOM		0 GLU	32	16.155	4.935	-3.992	1.00 28.96
ATOM	241		33	13.985		-4.140	
ATOM		CA GLU				-2.722	1.00 30.79
MOTA	243		33	12.483		-2.395	
ATOM	244	• •	33	12.198	6.452		1.00 47.15
ATOM .	245		33	10.798		-0.577	1.00 57.42
ATOM		OE1 GLU	33	9.828	6.400		1.00 63.55
ATOM	247	OE2 GLU			7.871	0.252	1.00 63.48
MOTA	248	C GLU	33	14.101	4.527	-1.971	1.00 28.96
ATON	249	O GLU	33	13.613	3.476	-2.391	1.00 28.97
ATOM.	250	N ALA	34	14.835	4.592	-0.864	•
ATOM	251	CA ALA		15.115		-0.069	1.00 29.99
MOTA	252	CB ALA	34	16.607.	3.314	0.234	1.00 26.15

FIG. 11A-6

MOTA	253	C	ALA	34	14.319	3.410	1.230	1.00 32.79
ATOM	254	0	ALA	34	14.272			1.00 32.79
ATOM	255	N	VAL	35	13.685		1.530	1.00 31.99
ATOM	256	CA	VAL	35	12.901		2.750	1.00 32.37
ATOM	257	CB	VAL	35	11.388		2.497	1.00 32.24
ATOM	258	CG1	VAL	35	11.132		1.902	1.00 32.24
ATOM	259	CG2	VAL	35	10.866	· · · · -	1.579	1.00 32.80
ATOM	260	-, C '	VAL	35	13.117			1.00 32.80
ATOM	261	0	VAL	35	13.609	· · · · 	2.564	1.00 33.29
ATOM	262	N	ALA	36	12.759	0.513	4.543	1.00 32.66
ATOM	263	CA	ALA	36	12.902		5.152	1.00 32.39
ATOM	264	CB /	ALA	36	13.444		6.577	1.00 30.92
ATOM	265	C	ALA	36	11.535		5_166_	1.00 32.53
ATOM	266	0	ALA	36	10.533		5.532	1.00 29.98
ATOM	267	N '	VAL	37	11.492	-2.720	4.749	1.00 34.45
ATOM	268	CA	VAL	37	10.240		4.729	1.00 37.01
ATOM	.269			37	9.919		3.316	1.00 39.07
ATOM	270	CG1		37	8.660	-4.841	3.352	1.00 41.91
MOTA	271	CG2		37	9.729	-2.810	2.366	1.00 40.40
ATOM	272			37	10.322	-4.629	5.690	1.00 37.16
ATOM	273			37	11.134	-5.534	5.514	1.00 36.61
ATOM	274			38	9.485	-4.592	6.720	1.00 37.96
ATOM	275			38	9.451	-5.655	7.713	1.00 39.49
ATOM	276			38	9.048	-5.086	9.077	1.00 38.70
ATOM	277		•	38	9.168	-6.066	10.236	1.00 38.05
ATOM	278			38	8.840	-5.378	11.554	1.00 40.91
ATOM	279			38	9.022	-6.309	12.737	1.00 46.69
ATOM	280			38	8.790	-5.598	14.026	1.00 49.50
-ATOM-	281			38	8.434	-6.688	7.246	1.00 40.71
ATOM	282			38	7.253	-6.379	7.084	1.00 40.05
ATOM	283			39	8.901	-7.910	7.016	1.00 42.81
ATOM	284			39	8.030	-8.983	6.553	1.00 45.99
ATOM	285			39	8.666	-9.730	5.364	1.00 45.59
ATOM	286				7.693	-10.765	4.818	1.00 46.73
ATOM	287	CG1		39	9.046	-8.728	4.270	1.00 44.50
ATOM	288	CD1		39	9.742	-9.349	3.075	1.00 49.55
ATOM	289			39	7.753	-9.977	7.675	1.00 48.22
ATOM	290			39	8.673	-10.593	8.210	1.00 48.95
ATOM	291			40			8.025	1.00 50.79
ATOM	292			40		-11.046	9.089	1.00 53.10
ATOM	293			40	5.604	-10.275	10.336	
ATOM	294	CG1 \	IAL 4	40	6.752	-9.471	10.927	1.00 55.17

FIG.11A-7

• • • •	MOTA	295	CG2	VAL	40	4.453	-9.352	9.963	1.00 49.52
	MOTA	296	С	VAL .	40	4.995	-12.016	8.656	1.00 55.18
	MOTA	297	0	VAL	40	3.925	-11.608	8.206	1.00 54.97
	ATOM .	298	N	ASP	41	5.277	-13.307	8.801	1.00 57.61
	ATOM	299	CA	ASP	41	4.327	-14.352	8.437	1.00 59.72
	MOTA	300	CB	ASP	41	5.077	-15.653	8.142	1.00 63.63
	MOTA	301	CG	ASP	41	4.183	-16.719	7.545	1.00 70.52
	ATOM	302	OD1	ASP	41 :	3.141	-17.036	8.157	1.00 69.83
	ATOM	303	: 0 D2	ASP	41	4.525	-17.244	6.465	1.00 74.90
	ATOM	304	C	ASP	41	3.352	-14.561	9.595	1.00 58.84
	ATOM	305	0	ASP	41	3.675	-15.233	10.575	1.00 57.65
	ATOM	306	N	MET	42	2.159	-13.984	9.477	1.00 59.02
	ATOM	307	CA	MET	42	1.142	-14.092	10.520	1.00 60.04
	ATOM	308	CB	MET :	42	100	-13.415		1.00 59.22
	MOTA	309	CG	MET	42	-0.036	-11.910	9.863	1.00 60.26
ď	ATOM	310	SD	MET	42	-1.552	-11.157	9.227	1.00 69.49
	ATOM	311	CE	MET	42	-2.295	-10.547	10.725	1.00 66.84
٠	MOTA	312	C	MET	42	0.847	-15.532	10.931	1.00 60.57
	ATOM	313	0	MET	42	0.297	-15.774	12.006	1.00 60.43
	ATOM	314	N	ALA	43	1.216	-16.483	10.078	1.00 61.75
	ATOM	315	CA	ALA	43	0.983	-17.898	10.358	1.00 63.27
•	ATOM	316	CB	ALA	43	0.675	-18.642	9.061	1.00 64.51
	MOTA	317	C	ALA	43	2.180	-18.538	11.054	1.00 63.20
	ATOM	318	.0	ALA	43	2.055	-19.596	11.672	1.00 64.09
	ATOM	319	N	ALA	44	3.337	-17.894	10.950	1.00 62.86
	MOTA	320	CA	ALA	44	4.555	-18.404	11.568	1.00 65.57
	ATOM	321	CB	ALA	44	5.777	-17.767	10.910	1.00 67.13
	ATOM	322	C	ALA	44	4.566	-18.135	13.071	1.00 69.55
	ATOM	323	0	ALA	44	5.527	-17.497	13.550	1.00 69.48
	MOTA	324	OT	ALA	44	3.614	-18.571	13.752	1.00 73.84
	ATOM	325	CB	CYS	48	1.032	-12.998	16.789	1.00 61.49
	ATOM	326	SG	CYS	48	-0.413	-12.709	17.840	1.00 66.53
	ATOM	327	C	CYS	48	-0.172	-12.208	14.752	1.00 58.42
	MOTA	328	0	CYS	48.	0.282	-11.074	14.587	1.00 58.49
	ATOM	329	N	CYS	48	1.950	-13.489	14.540	
	MOTA	330	CA	CYS	48	0.697	-13.320	15.332	1.00 59.76
	ATOM	331	N	PRO	49	-1.437	-12.524	14.431	1.00 57.61
	MOTA	332	CD	PRO .	49	-2.015	-13.880	14.439	
	MOTA	333	CA	PRO	49	-2.389		13.865	1.00 57.53
	ATOM	334	CB	PRO	49	-3.655		13.690	1.00 58.94
	ATOM	335	CG	PRO	49		•		1.00 60.96
	MOTA	336	C	PRO	49		-10.340		1.00 56.62

FIG.11A-8

	ATOM	337	0	PRO	49		-2.602	-9.205	14.273	1.00 56.98
	MOTA	338	N	GLU	50			-10.580	16.036	1.00 55.89
	MOTA	339	CA	GLU	50		-3.104	-9.502	16.985	1.00 54.74
	ATOM	340	CB	GLU	50			-10.072	18.306	1.00 57.44
	MOTA	341	CG	GLU	50		-3.950	-9.012	19.348	1.00 66.02
	MOTA	342	CD	GLU	50		-4.288	-9.606	20.701	1.00 00.02
	ATOM	343	0E1	GLU	50			-10.271	21.295	1.00 72.50
	ATOM	344	0E2	GLU	50	· ·	-5.428	-9.410	21.171	1.00 78.20
	ATOM	345	C	GLU	50		-1.846	-8.680	17.256	1.00 78.20
	ATOM	346	0	GLU	50		-1.846	-7.458	17.100	1.00 51.31
	ATOM	347	N	ALA	51		-0.779	-9.359	17.666	1.00 48.02
	MOTA	348	CA	ALA.	51		0.487	-8.701	17.969	1.00 45.05
	ATOM	349	CB	ALA	51		1.577	-9.747	18,180	1.00 42.33
	ATOM	350	C	ALA :	51		0.895	-7.734	16.862	1.00 44.76
	ATOM	351	0	ALA	51		1.156	-6.558	17.116	1.00 43.03
	ATOM :	•	N	ILE	52		0.940	-8.234	15.633	1.00 44.08
	ATOM	353	CA	ILE	52		1.318	-7.409	14.494	
	ATOM	354	CB	ILE	52		1.402	-8.275	13.199	1.00 43.74
	ATOM	355	CG2	ILE	52		0.009	-8.542	12.651	1.00 45.02
	ATOM	356	CG1	ILE	52	•	2.287	-7.588	12.154	1.00 46.00
	ATOM	357	CD1	ILE	52		1.728	-6.309	11.590	1.00 46.21
	ATOM	358	C	ILE	52		0.309	-6.267	14.321	1.00 39.96
	ATOM	359	0	ILE	52		0.686	-5.137	14.006	1.00 38.32
	ATOM 100	360		LYS	53		-0.968	-6.560	14.544	1.00 38.61
	ATOM		CA	LYS	53		-2.012	-5.548	14.412	1.00 38.67
	ATOM	362	CB	LYS	53		-3.394		14.612	1.00 40.27
	ATOM	363	CG	LYS	53		-4.205	-6.289	13.327	1.00 50.21
	ATOM	364	CD	LYS	53	٠.	-3.501	-7.151	12.289	1.00 54.17
	ATOM:	365		LYS	53		•4.213	-7.088	10.948	1.00 59.51
		366	NZ	LYS	53		-4.230	-5.702	10.405	1.00 57.13
	ATOM	367		LYS	53	٠.	-1.829			1.00 37.31
	ATOM	368	0	LYS	53		-2.105	-3.240	15.072	1.00 37.14
	MOTA	369 370	N	LYS	54		-1.370		16.602	1.00 35.35
٠	ATOM	371	CA CB	LYS LYS	54		1.155		17.612	1.00 32.72
	ATOM	372	CG	LYS	54 54			-4.332		1.00 32.05
	ATOM	373	CD	LYS	54 54		-0.850	-3.344	20.138	1.00 29.96
	ATOM	374	CE	LYS	54 54		-0.733	-4.081	21.465	1.00 31.32
	ATOM	375	NZ	LYS	54 54		-0.720	-3.119	22.644	1.00 32.19
	MOTA	376	C	LYS	54 54		0.527	-3.833	23.939	1.00 32.80
	ATON	370 377	0	LYS	54 54		0.070	-2.852	17.240	1.00 30.86
	ATOM	378		GLU	5 4 55		0.086	-1.636	17.432	1.00 29.26
	,, oa	5,0	14	GLU	22		1.092	-3.514	16.703	1.00 30.10

FIG.11A-9

	MOTA	379	CA	GLU	55	2.315	-2.832	16.299	1.00	30.95	5
	ATOM	380	CB	GLU	55	3.356	-3.838	15.791		28.04	
	MOTA	381	CG	GLU	55	4.719	-3.209	15.511		29.66	
	ATOM	382	CD	GLU	55	5.780	-4.224	15.120		30.54	
	· MOTA	383	0E1	GLU	55	5.708	-5.375	15.595		32.37	
	MOTA	384	0E2	GLU	55	6.699	-3.865	14.350		30.60	
	MOTA	385	С	GLU	55	2.004	-1.818	15.203		31.62	
	MOTA	386	0	GLU	55	2.552	-0.717	15.186		29.23	
	MOTA	387	N	ILE	56	1.121	-2.197			32.35	
	MOTA	388	CA	ILE	56	0.741	-1.300	13.203	1.00	31.04	1
	ATOM	389	CB	ILE	56	-0.135	-2.018	12.151	1.00	30.54	1
	MOTA	390	CG2	ILE	56	-0.659	-1.014	11.138	1.00	28.78	3
	MOTA		CG1		56	0.678	-3.108_	11.454	1.00	27.54	1
	MOTA	392	CD1	ILE	56	-0.104		10.397	1.00	28.92	2 -
	ATOM		C	ILE	56	-0.047	-0.134	13.785	1.00	29.35	5
	MOTA	394	0	ILE	56	0.185	1.022	13.432	1.00	26.97	7
		395	N	CYS	57	-0.974		14.686	1.00	28.75	5
4.5	ATOM	396	CA	CYS	57	-1.794	0.587	15.314	1.00	30.00	5
	MOTA	397	CB	CYS	57	-2.728	-0.030		1.00	100	
	MOTA	398		CYS	57	-3.764	1.186	17.224		42.92	
	ATOM	399	C	CYS	57	-0.907	1.630	15.986	* *** * *	27.0	
	MOTA	ta ta Tellin ta agrica	0	CYS	57	-1.043	2.825	15.742	N. T.	27.89	4.14
	ATOM	401	N	ILE	58	-0.001	1.166	16.838	1.00		
	ATOM			ILE	58	0.896	2.076		1.00		
	ATOM	403		***	58	1.810	1.305	18.522		29.7	
	ATOM	404		•	58	2.934	2.212		1.00		
	MOTA			ILE	58	0.968	0.787	19.691		28.0	
	ATOM	·		ILE .	<u>58</u>	1.773	0.086	20.780		29.2	
	ATOM ATOM	407		ILE	58	1.735		16.545		23.3	
	ATOM -	408		ILE	58	1.910	4.077	16.703		23.9	_
	ATOM	409		ASN	59	2.237	2.204	15.509		23.8	-
		410			59 50		2.882				
		411		ASN	59 50	3.547	1.873	13.461		28.5	
	ATOM	412	CG	ASN	59 50	4.951	1.372	13.764		31.9	
		413	•	ASN	59 50	5.929	2.102	13.598		32.0	
	ATOM	414		ASN	59 50		0.129			27.2	
•	ATOM	415	C	ASN	59 50	2.302	4.023			26.9	
	ATOM	416	0	ASN	59	2.900	5.045	13.457		24.7	
	·	417	N	LYS	60	0.999	3.856	13.595		28.9	
	ATOM	418	CA	LYS	60	0.207	4.892	12.936			
	ATOM	419	CB	LYS -	60		4.376	=		33.2	
	ATOM	420	CG	LYS	60	-1.254	3.289	11.574	1.00	39.3	1

FIG.11A-10

	ATOM	421	CD	LYS	60	-2.689	2.881	11.275	1.00 50.07
•	MOTA	422	CE	LYS	60	-2.751	1.811	10.199	1.00 63.36
	ATOM	423	NZ	LYS	60	-4.156	1.431	9.879	1.00 70.80
	MOTA	424	С	LYS	60	0.112	6.167	13.769	1.00 30.79
	MOTA	425	0	LYS	60	-0.255	7.225	13.261	1.00 32.02
	MOTA	426	N	MET	61	0.453	6.067	15.049	1.00 29.22
	MOTA	427	CA	MET	61	0.402	7.214	15.948	1.00 28.15
	MOTA	428	CB	MET	61	0.133	6.752	17.383	1.00 26.84
	MOTA	429	CG	MET	61	-1.123	5.934	17.601	1.00 33.92
	MOTA	430	SD	MET	61	-1.086	5.213	19.267	1.00 36.19
	MOTA	431	CE	MET	61	-1.338	6.689	20.282	1.00 35.78
	MOTA	432	C	MET	61	1.719	7.982	15.969	1.00 27.73
	MOTA	433	0	MET	61	1.773 _	9.126	16.419	1.00 30.14
	MOTA	434	N	LEU	62	2.772	7.346	15.474	1.00 26.12
	MOTA	435	CA	LEU	62	4.112	7.921	15.516	1.00 25.23
	ATOM	436	CB	LEU	62	5.129	6.786	15.574	1.00 24.11
	MOTA	437	CG	LEU	62	4.747	5.617	16.481	1.00 22.84
	ATOM	438	CD1	LEU	62	5.836	4.560	16.419	1.00 23.66
	MOTA	439	CD2	LEU	62	4.531	6.119	17.905	1.00 26.09
	MOTA	440	C	LEU	62	4.546	8.901	14.438	1.00 26.40
	MOTA	441	0	LEU	62	4.434	8.629	13.244	1.00 27.81
	MOTA	442	N	ASN	63	5.060	10.044	14.883	1.00 25.22
	MOTA	443	CA	ASN	63	5.576	11.064	13.981	1.00 24.06
	MOTA	444	CB	ASN	63	4.438	11.900	13.388	1.00 28.33
	MOTA	445	CG	ASN	63	4.938	12.925	12.399	1.00 31.22
	MOTA	446	OD1	ASN	63	5.933	12.696	11.711	1.00 34.87
	MOTA	447	ND2	2 ASN	63	4.249	14.058	12.310	1.00 31.84
	MOTA	448	C	ASN	• 63	6.564	11.961		1.00 21.00
	ATOM	449	0	ASN	63	6.202	13.010	15.240	1.00 20.80
	MOTA	450	N	HIS	. 64	7.818	11.525		1.00 20.38
	MOTA	451	CA	HIS	64	8.869	12.279	15.433	1.00 20.84
	MOTA	452	CB	HIS	64	8.896			1.00 20.13
	MOTA	453	CG	HIS	64	9.818			1.00 18.13
	MOTA	454	•	2 HIS	•	9.601		18.387	
	MOTA	455	ND:	I HIS	64	11.158	12.479	17.888	1.00 15.42
	ATOM	456		1 HIS	64	11.726	13.433	18.602	1.00 16.82
	ATOM	457	NEZ	2 HIS		10.804	14.324	18.917	
	MOTA	458	C	HIS	64	10.221	11.983	14.786	1.00 19.49
	MOTA	459	0	HIS	64	10.475	10.863	14.351	1.00 19.75
	MOTA	460	N	GLU	65	11.094	12.985	14.733	1.00 21.02
	MOTA	461	CA	GLU	65 .	12.397	12.816	14.100	1.00 21.68
	MOTA	462	CB	GLU	65	13.124	14.163	14.000	1.00 24.01

FIG.11A-11

							
MOTA	463	CG GLU	65	13.445	14.843	15.322	1.00 33.53
MOTA	464	CD GLU	65	12.284	15.643	15.885	1.00 41.84
MOTA	465	OE1 GLU	65	12.503	16.371	16.878	1.00 47.89
MOTA	466	OE2 GLU	65	11.158	15.548	15.346	1.00 41.02
MOTA	467	C GLU	65	13.323	11.781	14.733	1.00 22.44
ATOM	468	O GLU	65 ·	14.288	11.347	14.100	1.00 21.52
MOTA	469	N ASN	66	13.038	11.380	15.972	1.00 21.25
ATOM	470	CA ASN	66	13.873	10.383		1.00 20.43
ATOM	471	CB ASN	66	14.389	10.926	17.970	1.00 18.34
MOTA	472	CG ASN	66	15.360	12.089		2
MOTA	473	OD1 ASN	66	15.096	13.205	18.234	
MOTA	474	ND2 ASN	66	16.487			
MOTA	475	C ASN	66		9.055		1.00 20.35
ATOM	476	O ASN	66	13.463	8.278		1.00 18.70
MOTA	477	N VAL	67	12.146			1.00 19.53
ATOM	478	CA VAL	67	11.356			1.00 20.23
MOTA	479	CB VAL	67	9.935	7.840		
ATOM	480	CG1 VAL	67	9.074		16.470	1.00 17.62
ATOM	481	CG2 VAL	67	10.037		18.046	1.00 19.56
ATOM	482	C VAL	67	11.231		14.566	1.00 20.28
ATOM	483	O VAL	67	10.872		13.680	
ATOM	484	N VAL	68	11.541		14.333	
ATOM	485	CA VAL	68	11.449	5.247	12.991	1.00 20.39
MOTA	486	CB VAL	68	11.694	3.710	13.015	
MOTA	487	CG1 VAL	68	11.334	3.093	11.665	1.00 18.65
ATOM	488	CG2 VAL	68	13.155	3.420	13.327	1.00 16.32
MOTA		C VAL	68	10.074	5.542	12.393	1.00 22.68
MOTA	490	O VAL	68	9.046	5.217	12.986	1.00 22.91
ATOM		N LYS	69	10.068	6.172	11.221	1.00 24.55
ATOM	492	CA LYS	69	8.833		10.530	1.00 26.28
ATOM	493	CB LYS	69	9.129	7.465	9.353	1.00 31.62
ATOM	494	CG LYS	69	8.623	8.889	9.512	1.00 44.19
ATOM	495	CD LYS	69 🐰	9.589	9.741	10.314	1.00 51.62
MOTA		CE LYS	69	9.187	11.207	10.281	1.00 51.35
MOTA	497	NZ LYS	69	10.241	12.081	10.865	1.00 48.96
ATOM	498	C LYS	69	8.103	5.310	9.990	1.00 24.58
ATOM	499	0 LYS	69	8.729	4.348	9.539	1.00 25.04
ATOM	500	N PHE	70	6.776	5.368	10.040	1.00 25.47
ATOM	501	CA PHE	70	5.915	4.307	9.527	1.00 26.89
MOTA	502	CB PHE	70	4.824	3.961	10.545	1.00 29.09
MOTA	503	CG PHE	70	3.841	2.928	10.060	1.00 27.43
MOTA	504	CD1 PHE	70	4.248	1.621	9.808	1.00 28.02

FIG.11A-12

ATOM 506 CE1 PHE 70 3.337 0.659 9.372 1 ATOM 507 CE2 PHE 70 1.583 2.310 9.429 1 ATOM 508 CZ PHE 70 1.999 1.006 9.182 1 ATOM 509 C PHE 70 5.271 4.874 8.263 1	.00 30.32 .00 31.52 .00 29.79 .00 30.08 .00 28.21 .00 28.68 .00 29.81
ATOM 506 CE1 PHE 70 3.337 0.659 9.372 1 ATOM 507 CE2 PHE 70 1.583 2.310 9.429 1 ATOM 508 CZ PHE 70 1.999 1.006 9.182 1 ATOM 509 C PHE 70 5.271 4.874 8.263 1	.00 31.52 .00 29.79 .00 30.08 .00 28.21 .00 28.68
ATOM 507 CE2 PHE 70 1.583 2.310 9.429 1 ATOM 508 CZ PHE 70 1.999 1.006 9.182 1 ATOM 509 C PHE 70 5.271 4.874 8.263 1	.00 29.79 .00 30.08 .00 28.21 .00 28.68
ATOM 508 CZ PHE 70 1.999 1.006 9.182 1 ATOM 509 C PHE 70 5.271 4.874 8.263 1	.00 30.08 .00 28.21 .00 28.68
ATOM 509 C PHE 70 5.271 4.874 8.263 1.	.00 28.21 .00 28.68
ATOM FIG O DUE 70	.00 28.68
ATOM E11 N TWO 71	, UU ZY.OI
ATOM F12 CA TVD 71	.00 31.13
ATOM 512 CP TVD 71 F OF	.00 31.13
ATOM F14 CC TVD 71	.00 25.81
ATOM ETE ON TO	.00 28.78
ATOM E16 CE1 TVD 71	.00 23.78
ATOM E17 CD2 TVD 71	.00 26.54
ATOM E10 CE2 TVD 71 0 CO2	.00 24.23
ATOM 519 CZ TYR 71 9.763 6.442 5.209 1	00 23.46
ATOM _ 520 OH TYR 71 10.991 7.056 5.330 1	.00 29.32
ATOM 521 C TYR 71 3.634 4.049 5.520 1	.00 34.42
ATOM 522 0 TYR 71 2.842 4.596 4.753 1.	00 36.42
ATOM 523 N GLY 72 3.397 2.865 6.076 1.	00 34.34
ATOM 524 CA GLY 72 2.163 2.149 5.801 1.	00 33.61
ATOM 525 C GLY 72 2.392 0.653 5.757 1.	00 34.42
ATOM 526 0 GLY 72 3.511 0.191 5.972 1	00 34.69
ATOM 527 N HIS 73 1.341 -0.111 5.475 1.	00 37.74
ATOM 528 CA HIS 73 1.463 -1.564 5.413 1.	00 40.55
ATOM 529 CB HIS 73 1.102 -2.174 6.769 1.	00 39.94
ATOM 530 CG HIS 73 -0.340 -2.012 7.141 1.	00 41.03
AIOM 531 CD2 HIS 73 -1.017 -0.953 7.642 1.	00 38.25
ATOM 532 ND1 HIS 73 -1.265 -3.021 6.986 1.	00 42.56
ATUM 533 CET HIS 73 -2.452 -2.591 7.377 1.	00 39.22
AIOM 534 NE2 HIS 73 -2.329 -1.338 7.779 1.	00 37.48
ATUM 535 C HIS 73 0.576 -2.164 4.325 1.	00 42.07
ATUM 536 U HIS /3 -0.407 -1.553 3.907 1.	00 40.35
ATUM 53/ N ARG 74 0.933 -3.363 3.875 1.	00 45.26
ATOM 538 CA ARG 74 0.176 -4.056 2.837 1.	00 50.38
ATOM 539 CB ARG 74 1.022 -4.169 1.567 1.	
ATOM 540 CG ARG 74 1.382 -2.819 0.963 1.	00 63.87
ATUM 541 CD ARG 74 2.373 -2.946 -0.184 1.	00 70.66
ATUM 542 NE ARG /4 1.861 -3.752 -1.288 1.	00 72.42
ATOM 543 CZ ARG 74 2.485 -3.897 -2.453 1.	00 73.76
ATOM 544 NHI ARG 74 3.645 -3.289 -2.667 1.	00 64.85
ATOM 545 NH2 ARG 74 1.951 -4.650 -3.406 1.	00.78.24
AIDM. EAE C ADC 74 A AGE TO THE	00 52.53

FIG.11A-13

MOTA	547	O ARG	74	0.550	-6.237	3.785	1.00 50.95
MOTA	548	N ARG	75	-1.554	-5.725	3.148	1.00 56.77
MOTA	549	CA ARG	75	-2.138	-7.002	3.550	1.00 61.66
MOTA	550	CB ARG	75	-3.617	-7.046	3.150	1.00 66.26
MOTA	551	CG ARG	75 •	-4.406	-5.800	3.536	1.00 70.07
MOTA	552	CD ARG	75	-4.471	-5.610	5.043	1.00 75.18
ATOM	553	NE ARG	75	-5.229	-6.674	5.697	1.00 79.27
MOTA	554	CZ ARG	75	-5.442	-6.742	7.007	1.00 81.70
MOTA	555	NH1 ARG	75	-4.953	-5.806	7.810	1.00 80.67
MOTA	556	NH2 ARG	75	-6.147	-7.744	7.514	1.00 80.01
MOTA	557	C ARG	75	-1.404	-8.183	2.917	1.00 62.81
MOTA	558	O ARG	75	-0.570	-8.821	3.557	1.00 62.78
MOTA	559	N GLU	76	-1.730	411	and the second second	
MOTA	560	CA GLU	76	-1.109	-9.565	0.920	1.00 62.56
MOTA	561	CB GLU	76	0.399	-9.332	0.799	1.00 62.91
MOTA	562	CG GLU	76	1.081	-10.208	-0.240	
MOTA	563	CD GLU	76	0.711	-9.820	-1.659	1.00 71.31
MOTA	564	OE1 GLU	76	1.016	-8.676	-2.058	1.00 70.71
MOTA	565	OE2 GLU	76	0.116	-10.653		
MOTA	566	C GLU	76	-1.361	-10.931	1.561	
MOTA	567	O GLU	76	-0.420	-11.663	1.874	1.00 62.16
MOTA	568	N GLY	77	-2.632	-11.270	and the second second	and the state of t
MOTA	569	CA GLY	77		-12.551		(3)
MOTA	570	C GLY	77	-2.625	·12.690		and the second s
MOTA	571	O GLY			-12.078		
MOTA	572	N ASN	•		-13.501		
MOTA	573	CA ASN			-13.732		
MOTA	574				-15.236		
MOTA	575		=	• •	-15.914		
ATOM		OD1 ASN			-15.815	4.661	· ·
MOTA	577	ND2 ASN	78		-16.613		•
MOTA	•	C ASN			-13.096		
MOTA	579	0 ASN			-13.079		
ATOM	.580	N ILE	79	0.831	-12.579		· ·
MOTA	581	CA ILE			-11.947		·
ATOM	582	CB ILE		•	-12.281		
ATOM	583	CG2 ILE			-11.570	3.864	
MOTA	584	CG1 ILE			-13.795	3.607	1.00 52.83
ATOM	585	CD1 ILE			-14.415	4.846	1.00 50.48
MOTA	586		79		-10.433	4.991	1.00 52.67
MOTA	587	0 ILE	: 79	1.668	-9.767		
MOTA	588	'N GLN	80	2.272	-9.895	6.180	1.00 50.66

FIG.11A-14

MOTA	589			2.191	-8.455	6.408	1.00 48.22
MOTA	590	CB GLA		1.944	-8.158	7.891	1.00 50.12
MOTA	591	CG GLN		0.521	-8.399	8.362	1.00 48.27
MOTA	592	CD GLN	80	-0.494	-7.572	7.594	1.00 49.09
MOTA	593	OE1 GLA	80	-0.372	-6.351	7.493	1.00 45.08
MOTA	594	NE2 GLM	1 80	-1.506	-8.237	7.049	1.00 58.25
MOTA	595	C GLM	80	3.469	-7.750	5.966	1.00 46.07
MOTA	596	O GLA	80	4.572	-8.222	6.238	1.00 45.36
MOTA	597	N TYP	81	3.307	-6.618	5.288	1.00 45.11
MOTA	598	CA TYP	81	4.436	-5.829	4.805	1.00 43.61
MOTA	599 .	CB TYP	81	4.385	-5.706	3.280	1.00 42.41
MOTA	600	CG TYP	81	4.641	-7.001	2.545	1.00 43.09
MOTA	601	CD1 TYP	81	5.918	-7.559	2.504	1.00 40.63
MOTA	602	CE1 TYP	81	6.157	-8.756	1.834	1.00 45.03
MOTA	603	CD2 TYP	81	3.606	-7.672	1.896	1.00 41.43
ATOM	604	CE2 TYP	₹ 81	3.835	-8.870	1.225	1.00 44.97
MOTA	605	CZ TYP	₹ 81	5.111	-9.405	1.197	1.00 46.87
MOTA	606	OH TYP	₹ 81	5.339	-10.589	0.534	1.00 49.59
MOTA	607	C TY	R 81	4.409	-4.433	5.419	1.00 42.40
MOTA	608	0 TY		3.585	-3.602	5.042	1.00 43.43
MOTA	609	N LEI		5.309	-4.178	6.365	1.00 39.70
ATOM	610	CA LE		5.372	-2.874	7.010	1.00 37.19
MOTA	611	CB LEI		5.616	-3.028	8.517	1.00 39.36
ATOM	612	CG LE		4.579	-3.785	9.358	1.00 38.76
ATOM	613	CD1 LE		4.968		10.827	1.00 32.68
MOTA	614	CD2 LE				9.155	1.00 39.92
ATOM	615	C LE		6.485		6.397	1.00 34.17
ATOM	616	0 LEI		7.659	-	6.445	1.00 32.38
ATOM	617	N PH					1.00 33.55
ATOM		CA PH		7.083	0.008	5.209	1.00 31.38
ATOM	619	CB PH		6.464	- · · · · · · · · · · · · · · · · · · ·	4.011	1.00 36.90
ATOM	620	CG PH					1.00 40.96
ATOM	621						1.00 42.10
ATOM	622	CD2 PH			•		1.00 41.38
ATOM	623	CE1 PH					1.00 41.93
ATOM	624	CE2 PH					1.00 40.00
ATOM	625	•	E 83				1.00 39.51
ATOM	626	C PH				6.251	· · · -
ATOM	627	O PH		· · · · · · · · · · · · · · · · · · ·			1.00 24.97
ATOM	628	N LE					1.00 27.73
ATOM	629	CA LE					1.00 27.95
ATOM	630	CB LE	J 84	9.837	0.837	8.84 6.	1.00 28.57

FIG.11-A15

MOTA	631	CG	LEU	84	8.720	-0.038	9.430	1.00 24.56
MOTA		CD1		84	9.313	-1.254	10.122	1.00 21.64
MOTA	633	CD2	LEU	84	7.874	0.787	10.386	1.00 24.46
MOTA	634	C	LEU	. 84	10.604	2.508	7.164	1.00 28.88
MOTA	635	0	LEU	84	11.204	2.184		1.00 28.67
MOTA	636		GLU	85	10.949	3.551		1.00 28.76
MOTA	637		GLU	85	12.072	4.404		1.00 28.20
MOTA	638	CB	GLU	85	12.170		•	1.00 29.51
MOTA	639	CG	GLU	85 .	13.371	6.450	•	1.00 34.24
ATOM	640		GLU	85	13.405	7.556		1.00 36.83
MOTA	641	0E1		85	14.354	8.367		1.00 36.87
MOTA	642	0E2	GLU	85	12.478		10.280	
ATOM	643	С	GLU	⁻ 85	13.367		•	1.00 27.67
ATOM 4	644	0	GLU	85	13.645			1.00 28.74
MOTA	645	N	TYR	86	14.150		the second secon	1.00 24.64
MOTA	646	CA	TYR	86	15.421	3.059		1.00 23.98
ATOM	647	CB	TYR	86	15.793	2.870	4.901	1.00 24.65
ATOM	648	CG	TYR	86	17.208			1.00 25.72
ATOM	649	CD1	TYR	86	17.652			1.00 22.83
ATOM	650	CE1	TYR	86	18.954	0.719	5.014	1.00 20.99
ATOM	651	CD2	TYR	86	18.103	3.108	3.888	1.00 24.91
MOTA	652	CE2	TYR	86	19.404	2.659	3.668	1.00 25.43
MOTA	653	CZ	TYR	86	19.822	1.466	4.234	1.00 24.00
MOTA	654	OH	TYR	86	21.110	1.021	4.030	1.00 31.37
- ATOM -	655	. C .	TYR	86	16.499	3.879	7.081	1.00 24.51
ATOM -	656	0	TYR	86	16.544	5.075	6.828	1.00 25.53
MOTA	657	N	CYS	87	17.241	3.236	7.974	1.00 23.81
MOTA	658	CA	CYS	87	18.303	3.914		1.00 24.33
MOTA	659	CB	CYS	87	18.059	3.747	10.218	1.00 21.77
ATOM		SG	CYS	87	16.439	4.374		1.00 22.54
MOTA	661	C	CYS		19.637	3.310		1.00 22.59
ATOM	662	. 0		- 87	20.090	2.310	8.840	1.00 22.98
MOTA	663		SER		20.263	3.935	7.291	1.00 22.63
MOTA	664	CA	SER	. ••	21.519	3.452	6.729	1.00 23.13
MOTA	665		SER	88	21.869	4.258	5.470	1.00 23.11
ATOM	666	0G	SER		22.008	5.641	5.750	1.00 27.21
MOTA	667	€ į	SER	88	. 22.727	3.412	7.656	1.00 24.35
MOTA	. 668	Ó	SER	88	23.746	2.808	7.318	1.00 26.40
ATOM	669	N	GLY	89	22.618	4.049	8.818	1.00 23.40
MOTA	670	CA	GLY	89 .	23.720	4.053	9.764	1.00 21.46
MOTA	671	C	GLY	89	23.777	2.793	10.613	1.00 21.62
MOTA	672	0 -	GLY	· 89	24.747	2.566	11.336	1.00 23.85

FIG.11A-16

				•					
MOTA	673	N	GLY	90	••	22.736	1.974	10.523	1.00 19.99
MOTA	674	CA	GLY	90		22.700		11.275	1.00 19.98
MOTA	675	С	GLY	90		22.263	0.895	12.723	1.00 19.03
ATOM	676	0	GLY	90		21.563	1.845	13.066	1.00 19.39
ATOM	677	N	GLU	91		22.689	-0.036	13.569	1.00 20.36
ATOM	678	CA	GLU	91		22.325	-0.017	14.983	1.00 20.23
MOTA	679	CB	GLU	91		22.202	-1.439	15.522	1.00 21.22
ATOM	680	CG	GLU	91		21.218	-2.329	14.792	1.00 23.61
ATOM	681	CD	GLU	91 .		21.215	-3.743	15.342	1.00 23.88
ATOM	682	0E1	GLU	91		20.492	-4.594	14.784	1.00 29.99
MOTA	683	0E2	GLU	91		21.934	-4.000	16.334	1.00 22.19
MOTA	684	С	GLU	91		23.334	0.721	15.846	1.00 20.72
ATOM	685	0	GLU	91		24.526	0.739	15.556	1.00 20.19
MOTA	686	N	LEU	92		22.847	1.311	16.932	1.00 19.98
ATOM	687	CA	LEU	92		23.712	2.020	17.864	1.00 19.39
ATOM	688		LEU	92	•	22.868	2.671	18.963	1.00 18.03
MOTA	689	CG	LEU	92		23.616	3.333	20.122	1.00 19.18
MOTA	690		LEU	92		24.427	4.513	19.612	1.00 22.62
ATOM	691		LEU	92		22.596	3.783	21.176	1.00 16.79
MOTA	692	C	LEU	92		24.641	0.989	18.480	1.00 19.97
MOTA	693	0	LEU	92		25.781	1.284	18.834	1.00 19.58
MOTA	694	N	PHE	93		24.134	-0.232	18.599	1.00 20.27
ATOM	695	CA	PHE	93		24.895	-1.322	19.178	1.00 21.88
ATOM	696		PHE	. 93		24.099	-2.628	19.058	1.00 26.11
ATOM	697	CG	PHE	93		24.813	-3.834	19.611	1.00 28.84
MOTA	698	CD1.		93		25.734	-4.533	18.836	1.00 29.47
ATOM	699		PHE	93		24.561	-4.274	20.907	1.00 31.53
ATOM	700	CE1		93		26.393	-5.656	19.344	1.00 30.19
MOTA	701		PHE	93		25.216	-5.397	21.425	1.00 36.41
ATOM	702		PHE	93		26.132	-6.088	20.641	1.00 30.51
MOTA	703	C	PHE	93	•	26.245	-1.458	18.481	1.00 21.58
ATOM	704		PHE	93		• •	-1.675		1.00 20.32
MOTA	705		ASP			26.245	-1.300	17.161	1.00 23.68
MOTA	706		ASP			27.474			1.00 25.61
ATOM	707	CB	ASP	94	•	27.118	-1.757	14.925	1.00 30.05
ATOM	708		ASP	94	•	26.495	•	14.782	1.00 31.43
ATOM	709	OD1		94		25.725	-3.361	13.827	1.00 33.18
ATOM	710	0D2		94		26.783	-4.011	15.628	1.00 34.06
MOTA	.711		ASP	94		28.423	-0.232	16.451	1.00 24.95
MOTA	712	0	ASP	94		29.501	-0.257	15.860	1.00 27.73
ATOM	713	N	ARG	95		28.035		17.194	1.00 23.99
MOTA	714	CA	ARG	95		28.870	1.991	17.363	1.00 23.16

FIG.11A-17

MOTA	715	CB A	NRG	95	28.008	3.255	17.263	1.00 24.65
MOTA	716	CG A	ARG	.95	27.399	3.479	15.888	1.00 29.91
MOTA	717	CD A	ARG	95	28.488	3.806	14.875	1.00 39.68
ATOM	718	NE A	ARG	95	29.148	5.055	15.241	1.00 47.46
ATOM -	719	CZ /	ARG	95	28.687	6.262	14.929	1.00 46.44
ATOM	720	NH1	ARG	95	27.568	6.386	14.227	1.00 39.38
ATOM	721	NH2	ARG	95	29.325	7.346	15.353	1.00 42.03
ATOM	722	C /	ARG	95	29.557	1.935	18.727	1.00 22.01
MOTA	723	0 /	ARG	95	30.340	2.819	19.090	1.00 21.05
MOTA	724	N :	ILE	96	29.246	0.885	19.482	1.00 22.78
ATOM	725	CA :	ILE	96	29.811	0.680	20.806	1.00 22.43
ATOM	726	CB :	ILE	96	28.735	0.146	21.776	1.00 20.79
MOTA	727	CG2	ILE	96	29.332	-0.066	23.160	1.00 20.67
MOTA	728	CG1		96	27.578	1.146	21.845	1.00 18.29
ATOM	729	CD1	ILE	96	26.357	0.640	22.590	1.00 18.91
MOTA		C	ILE.	96	30.963	-0.315	20.732	
MOTA	731		ILE	96	30.769	-1.469	20.360	1.00 25.11
MOTA	732		GLU	97	32.162	0.136	21.076	
MOTA	733		GLU	97	33.318		21.038	
ATOM	734		GLU	97	34.605	0.055	20.849	
MOTA	735		GLU	97	34.830	er and the second second	19.444	ing the contract of the contract of the contract of
MOTA	736	erit a e de la comi	GLU	97	33.846	1.719		1.00 53.97
MOTA	737	0E1		97	33.759	2.740		
ATOM	738	0E2		97	33.165			1.00 58.08
ATOM			GLU	97	33.383	-1.570	22.325	1.00 24.68
MOTA	740		GLU	97	33.415			1.00 23.06
ATOM	741		PRO	98	33.395	-2.906		1.00 24.21
ATOM	742		PRO	98	33.233	-3.720	20.987	
MOTA	743		PRO	98			23.392	1.00 23.90
MOTA	744	0	PRO	98	33.695		22.792	1.00 23.15
· · · · · · · · · · · · · · · · · · ·	745		PRO	98	32.877	-5.090	21.560	1.00 21.24
	746		4.4	98			A CONTRACTOR OF THE CONTRACTOR	1.00 24.35
ATOM			PRO	98				1.00 25.72
ATOM		N	** *	•	34.071			•
ATOM		CA	** *	•	34.919	*		•
ATOM	750		ASP	99	36.119			
ATOM	751		ASP	99	35.726		27.365	
MOTA	752			99	35.109		26.579	
ATOM	_	OD2		99	36.030		28.531	
MOTA	754		ASP	99			26.826	•
ATOM	755		ASP	99			27.741	
ATOM	<i>7</i> 56	N	ILE	100	35.066	-0.662	25.841	1.00 23.29

FIG.11-A18

MOTA	757	CA	ILE	100		35.532	0.721	25.862	1.00 23.28
ATOM	758	CB	ILE	100		36.625	1.000	24.786	1.00 28.85
ATOM	759	CG2		100		37.699	-0.076	24.842	1.00 30.41
ATOM	760	CG1		100		36.017	1.042	23.393	1.00 37.12
MOTA	761	CD1	ILE	100		37.017	1.438	22.311	1.00 50.18
ATOM	762	C	ILE	100		34.403	1.737	25.699	1.00 20.45
ATOM	763	0	ILE	100		34.413	2.771	26.354	1.00 20.39
MOTA	764	N	GLY	101		33.447	1.445	24.823	
MOTA	765	CA	GLY	101		32.334	2.355	24.610	1.00 19.62
ATOM	766	C	GLY	101		32.521	3.227	23.384	
ATOM	. 767	0	GLY	101		32.745	2.721	22.285	1.00 19.10
ATOM	768	N	MET	102		32.410	4.539	23.570	1.00 18.46
ATOM	769	CA	MET	102		32.583			1.00 17.81
ATOM	770	CB	MET	102		31.291	5.655	21.676	1.00 19.03
ATOM	771	CG	MET	102		30.170	6.358	22.449	1.00 19.18
MOTA	772	SD	MET	102		28.677	6.592	21.435	1.00 16.18
ATOM	773	CE	MET	102		28.107	4.874	21.273	1.00 15.36
MOTA	774	C	MET	102		32.931	6.853	23.120	1.00 18.77
ATOM	77 5	0.	MET	102		32.784	7.028	24.331	1.00 19.16
ATOM	776	Ň	PRO	103		33.403	7.821	22.317	1.00 18.19
ATOM	777	CD	PRO	103		33.736	7.749	20.882	1.00 16.39
MOTA	778	CA	PRO	103		33.749	9.138	22.863	1.00 18.51
ATOM	779	CB	PRO	103		34.109	9.940	21.619	1.00 17.03
ATOM	780		PRO	103		34.696	8.903	20.725	1.00 15.88
MOTA	781	С	PRO	103		32.562	9.741	23.614	1.00 19.83
MOTA	782	. 0	PRO	103		31.437	9.710	23.126	1.00 19.36
ATOM	783		GLU	104		32.823	10.290	24.794	1.00 18.81
ATOM	784	• .	GLU	104		31.771	10.873	25.617	1.00 19.36
MOTA	785		GLU	104	•	32.386	11.511	26.864	1.00 17.66
MOTA	786		GLU	104		31.406	11.735	27.996	1.00 16.18
MOTA	787		GLU	104	<i>i</i> .	32.058	12.320	29.231	1.00 21.03
ATOM				104		31.679	13.448	29.619	1.00 21.27
ATOM	_	0E2		104		32.946	11.657	29.819	1.00 19.74
MOTA		C		104		30.871	11.880	24.898	1.00 19.92
MOTA	791		GLU	104		29.653	11.886	25.105	1.00 19.71
ATOM			PRO ·	105	•	31.448	12.748	24.049	1.00 20.08
ATOM	793		PRO ·			32.877	13.000	23.789	1.00 20.80
MOTA	794		PRO-	105		30.607	13.723	23.342	1.00 16.33
ATOM	795			105		31.621	14.529	22.530	1.00 16.90
ATOM	· 796		PRO	105		32.875	14.459	23.403	1.00 16.88
MOTA	797		PRO	105		29.572	13.017	22.452	1.00 16.27
MOTA	· 798	0	PRO	105		28.424	13.452	22.344	1.00 17.44

FIG.11A-19

ATOM	799	N ASP	106	29.995	11.934	21.809	1.00 15.51	
MOTA	800	CA ASP	106	29.119	11.153	20.938	1.00 17.98	
MOTA	801	CB ASP	106	29.906	10.029	20.264	1.00 20.63	
MOTA	802	CG ASP	106	30.890	10.530	19.224	1.00 25.04	
MOTA	803	OD1 ASP	106	31.277	11.712	19.273	1.00 27.36	
MOTA	804	OD2 ASP	106	31.290	9.721	18.364	1.00 31.22	٠
MOTA	805	C ASP	106	28.001	10.522	21.771	1.00 15.08	. (7
MOTA	806	O ASP	106	26.829	10.515	21.375	1.00 16.13	105
MOTA	807	N ALA	107	28.371	9.980	22.925	1.00 14.46	. 2 - 1
ATOM	808	CA ALA	107	27.392	9.348	23.802	1.00 16.06	4
ATOM .	809	CB ALA	107	28.095	8.697	24.989	1.00 14.46	
ATOM	810	C ALA	107	26.363	10.373	24.288	1.00 15.73	is to
ATOM	8T1	O ALA	107	25.163	10.077	24.372	1.00 15.16	875
ATOM	812	N GLN	108	26.828	11.577	24.603	1.00 14.18	
ATOM	813	CA GLN	108	25.932	12.630	25.075	1.00 14.33	1/2
MOTA	814	CB GLN	108	26.722	13.874	25.492	1.00 17.52	1 gh
MOTA	815	CG GLN	108	25.868	14.876	26.277	1.00 16.31	723
ATOM	816	CD GLN	108	26.454			1.00 18.32	
MOTA	817	OE1 GLN	108	26.514	16.924		1.00 20.27	
ATOM	818	NE2 GLN	108	26.859	The second secon		1.00 12.37	
ATOM	819	C GLN	108	24.927	13.029		1.00 15.75	
ATOM	820	O GLN	108	23.745	13.212		1.00 14.23	
MOTA	821	N ARG	109	25.402	13.185			
ATOM	822	CA ARG	109	24.526		21.649		
ATOM	823	CB ARG	109	25.356			1.00 13.79	
MOTA	824		109	24.552		19.160	1.00 17.49	4.5
ATOM	825	CD ARG	109	25.408	2	17.902	1.00 23.46	
MOTA	826	NE ARG	109	25.536	12.928			
ATOM	827	CZ ARG	109	24.873		16.294		
ATOM	828	NH1 ARG	109	24.035		15.636		
		NH2 ARG	109	25.034		15.910	i i	
	830		109	23.473		-		
ATOM	831	O ARG	109	22.285		21.243		
MOTA	: 832		110	23.904		21.424		
ATOM	833		110	22.963	10.099			
ATOM	834					21.151		
MOTA	835	•	110	24.421		19.868		
ATOM	836		110	23.818			1.00 18.08	
ATOM	837		110	25.714		· ·		
ATOM	838		110	24.502		17.436		
ATOM	839		110	26.401				
MOTA	840	CZ PHE	110	25.798	7.991	17.481	1.00 14.35	

FIG.11A-20

MOTA	841	C PHE	110	21.962	10.059	22.366	1.00 12.75
ATOM	842	O PHE	110	20.777	9.777	22.155	1.00 12.75
MOTA	843	N PHE	111	22.435	10.339	23.579	1.00 13.13
MOTA	844	CA PHE	. 111	21.554	10.325	24.743	1.00 12.28
MOTA	845	CB PHE	111	22.367	10.450	26.039	1.00 15.94
ATOM	846	CG PHE	111	21.565	10.174	27.273	1.00 13.94
MOTA	847	CD1 PHE	111	21.146	8.877	27.566	1.00 13.02
MOTA	848	CD2 PHE	111	21.183	11.212	28.119	1.00 12.96
MOTA	849	CE1 PHE	111	20.354	8.617	28.683	1.00 10.93
ATOM	850	CE2 PHE	111	20.391	10.969	29.239	1.00 10.93
ATOM	851	CZ PHE	111	19.971	9.662	29.523	1.00 9.75
MOTA	852	C PHE	111	20.519	11.454		1.00 12.76
MOTA	853	O PHE	111	19.366	11.278		1.00 12.70
ATOM	854	N HIS	112	20.938	12.608	24.144	1.00 13.86
MOTA	855	CA HIS	112	20.027	13.742	23.970	1.00 14.60
MOTA	856	CB HIS	112	20.760	14.924	23.331	1.00 15.24
MOTA	857	CG HIS	112	21.699	15.642	24.249	1.00 15.45
MOTA	858	CD2 HIS	112	21.734	15.739	25.599	1.00 17.90
MOTA	859	ND1 HIS	112	22.718	16.444	23.779	1.00 17.44
MOTA	860	CE1 HIS	112	23.336	17.009	24.802	1.00 15.77
MOTA	861	NE2 HIS	112	22.757	16.598	25.918	1.00 22.23
ATOM	862	C HIS	112	18.903	13.339	23.019	1.00 15.61
MOTA	863	O HIS	112	17.726	13.619	23.263	1.00 16.06
MOTA	864	N GLN	113	19.276	12.699	21.915	1.00 14.44
MOTA	865	CA GLN	113	18.294	12.283	20.925	1.00 14.83
MOTA	866	CB GLN	113	18.998	11.869	19.635	1.00 12.67
MOTA	867	CG GLN	113	19.743	13.047	19.012	1.00 12.96
MOTA	868	CD GLN	113	20.508	12.662	17.764	1.00 22.01
MOTA	869	OE1 GLN	113	20.468	11.514	17.327	1.00 25.13
ATOM	870	NE2 GLN	113	21.218	13.625	17.186	1.00 21.52
MOTA	871	C GLN	113	17.406	11.170	21.450	1.00 14.34
ATOM	872			16.218	11.124	21.140	1.00 13.52
MOTA	873	N LEU	114	17.970	10.294	22.273	1.00 14.36
MOTA	874	CA LEU		17.177	9.217	22.863	1.00 13.96
MOTA	875	CB LEU	114	18.075	8.287	23.683	1.00 13.60
MOTA	876	CG LEU	114	17.404	7.167	24.485	1.00 14.47
MOTA	877		114	16.559	6.292	23.575	1.00 12.86
MOTA	878	CD2 LEU	114	18.491	6.320	25.175	1.00 11.34
ATOM	879	C LEU	114	16.109	9.848	23.775	1.00 13.12
MOTA	880	O LEU	114	14.925	9.483	23.730	1.00 12.42
MOTA	881	N MET	115	16.521	10.806	24.597	1.00 13.13
MOTA	882	CA MET	115	15.568	11.476	25.486	1.00 14.41

FIG.11A-21

ATOM	883	СВ	MET	115	16.274	12.516	26.367	1.00 13.67
ATOM	884	CG	MET	115	17.130		27.481	1.00 16.47
ATOM	885	SD	MET	115	16.170	10.931	28.639	1.00 16.82
MOTA	886	CE	MET	115	16.565	9.273	27.955	1.00 11.48
ATOM	887	C	MET	115	14.467	12.175	24.685	1.00 14.97
MOTA	888	0	MET	115	13.297	12.136	25.059	1.00 15.73
MOTA	889	N	ALA	116	14.842	12.819	23.585	1.00 16.10
ATOM	890	CA	ALA	116	13.859		22.752	1.00 15.36
ATOM	891	CB	ALA	116	14.551	14.203	21.581	1.00 14.43
MOTA		C	ALA	116	12.818		22.244	1.00 15.44
ATOM	893	0	ALA	116	11.617	12.785	22.269	1.00 17.74
ATOM	894	N	GLY	117	13.286	11.342	21.815	1.00 13.85
MOTA	895		GLY	117	12.379	10.322	21.312	1.00 13.18
MOTA	896	C	GLY	117	11.490	9.760	22.406	1.00 13.23
MOTA	897	0	GLY	117	10.294	9.563	22.204	1.00 15.51
ATOM	898	N	VAL	118	12.068	9.484	23.571	1.00 13.22
ATOM	899	CA	VAL	118	11.275	8.944	24.669	
ATOM	900	CB	VAL	118	12.184	8.412	25.790	
MOTA	901	CG1	VAL	118	11.343	7.955	26.981	
ATOM	902	CG2	VAL	118	12.999	7.250	25.256	
ATOM	903	C	VAL	118	10.277			and the state of t
ATOM	904	0	VAL	118	9.150	9.598		and the second s
ATOM	905	N	VAL	119		11.230		1.00 15.14
MOTA	906	CA	VAL	119	9.764		25.809	
MOTA	907	CB	VAL	119	10.428	13.682	25.807	1.00 15.22
MOTA	908		L VAL	119				1.00 16.36
MOTA	909	CG2	2 VAL	119				1.00 10.99
ATOM	910	<u> </u>	VAL	119		and the second section of the second section	24.889	1.00 15.40
ATOM		0		119	7.401		25.341	1.00 16.23
ATOM -	912			120			23.596	1.00 13.60
ATOM	913	CA	TYR	120	7.737		22.610	1.00 15.80
ATOM	914	· CB	TYR				21.201	
ATOM	915	CG	TYR		7.266		20.151	1.00 16.47
ATOM			1 TYR	•	6.407	12.969		
ATOM	917	CE:	1 TYR	120	5.373	,	18.861	1.00 16.07
ATOM	918	CD	2 TYR	120	7.080		19.593	•
ATOM			2 TYR	120	6.055			
ATOM	920	CZ		120	5.205			
MOTA	921	OH	TYR	120	4.176	11.243		
ATOM	922	C	TYR	120	6.774		22.818	1.00 17.10
ATOM	923		TYR	120	5.553	•		1.00 15.76
MOTA	924	N	LEU	121	7.320	9.847	22.880	1.00 14.94 ·

FIG.11A-22

MOTA	925	CA LEU	121	6.	491	8.670	23.074	1.00	14.73
MOTA	926	CB LEU	121	.7.	351	7.407	23.136	1.00	14.27
ATOM	927	CG LEU	121	. 8.	129	7.046	21.867	1.00	15.46
MOTA	928	CD1 LEU	121	8.	970	5.796	22.125	1.00	15.26
MOTA	929	CD2 LEU	121	· 7.	165	6.825	20.711	1.00	11.93
MOTA	930	C LEU	121	5.	661	8.782	24.346	-	16.08
MOTA	931	0 LEU	121		453	8.565	24.328		14.61
MOTA	932	N HIS			309	9.126	25.452		14.83
ATOM	933	CA HIS	122		594	9.234	26.710		15.63
MOTA	934	CB HIS	122		585	9.508	27.842		15.63
MOTA	935	CG HIS			434	8.321	28.179		12.77
ATOM	936	CD2 HIS			432	7.061	27.686		12.46
ATOM	937	ND1 HIS		•	402		.:29.160		12.14
MOTA	938	CE1 HIS			957		29.260		11.43
MOTA	939	NE2 HIS			385	6.352	28.377		12.08
MOTA	940	C HIS			515	10.307			17.17
ATOM	941	0 HIS		· -	452	10.163	27.246		17.14
MOTA	942	N GLY			.783	11.362	25.886		17.18
MOTA	943	CA GLY			.818	12.439	25.755		19.68
MOTA	944	C GLY			.536	12.000	25.071		19.70
MOTA	945	0 GLY	123		.468	12.559	25.333		22.02
MOTA	946	N ILE			.636	11.006	24.195		18.52
ATOM	947	CA ILE			.468	10.489	23.485		20.43
MOTA	948	CB ILE	124	1.	807	10.171	21.992		26.21
MOTA	949	CG2 ILE	124	2.	.795	9.024	21.896	1.00	22.32
MOTA	950	CG1 ILE	124	. 0.	.531	9.825	21.219	1.00	43.02
ATOM	951	CD1 ILE	124	-0.	.447	10.975	21.093	1.00	56.77
ATOM	952	C ILE	124	0.	.922	9.246	24.200	1.00	19.54
ATOM	953	0 ILE	124	0	.023	8.569	23.705	1.00	20.58
ATOM	954	N GLY		1.	.468	8.953	25.379	1.00	19.44
ATOM	955	CA GLY	125	· . 0	.989	7.816	26.148	1.00	17.08
MOTA	956	C GLY		1.	.490	6.438	25.753	1.00	15.22
ATOM	957	0 GLY		. 0	.872	5.425	26.100	1.00	15.72
ATOM	958	N ILE		2	.593	6.368	25.022	1.00	14.59
ATOM	959	CA ILE		3	.098	5.054	24.669	1.00	15.92
ATOM	960	CB ILE		3	.197	4.831	23.121	1.00	23.85
MOTA	961	CG2 ILE		1	. 985	5.439	22.415	1.00	21.96
MOTA	962	CG1 ILE		4.	.478	5.425	22.565	1.00	25.44
MOTA	963	CD1 ILE		4	.761	4.944	21.151	1,00	32.08
ATOM	964	C ILE		. 4	.452	4.759	25.304	1.00	14.58
ATOM	965	0 ILE		. 5	.301	5.645	25.466	1.00	13.21
MOTA	966	N THR	127	4	.619	3.513	25.725	1.00	15.33

FIG.11A-23

MOTA	967	CA	THR	127	** ****	5.884	3.077	26.301	1.00	16.98	
MOTA	968		THR	127		5.710	2.492	27.730		21.08	••
MOTA	969	OG1	THR	127	: .	6.963	1.951	28.171		42.01	
MOTA	970	CG2	THR	127		4.657	1.398	27.753	1.00	8.51	
MOTA	971		THR	127		6.458	2.024	25.350	1.00	15.46	
ATOM	972	0	THR	127	***	5.738		24.862		13.84	
ATOM	973	N	HIS	128		7.757		25.084		16.45	, · .
ATOM	974	CA	HIS	128		8.415		24.152		14.14	
MOTA	975	CB	HIS	128		9.736		23.696		16.06	
MOTA	976	CG	HIS	128		10.479		22.693		19.22	
MOTA	977	CD2	HIS	128		10.596		21.349		20.26	
MOTA	978	ND1	HIS	128		11.214		23.043			.11.
ATOM	979	CE1	HIS	128		11.754		21.958		19.24	127
MOTA	980	NE2	HIS	128		11.394	0.082	20.916		19.76	
MOTA	981	С	HIS	128		8.635	-0.199	24.755		13.34	
ATOM	982	0	HIS	128	, Same	8.422	-1.219	24.087	1.00	13.46	444
MOTA	983	N	ARG	129		9.044	-0.215	26.025	1.00	12.33	-3.55
MOTA	984	CA	ARG	129		9.283	-1.427	26.820	1.00	13.03	n With
MOTA	985	-CB	ARG	129		7.998	-2.267	26.897	1.00	11.47	
MOTA	986	CG	ARG	129	() 4- Jul	6.825	-1.460	27.467	1.00	16.16	
MOTA	987	CD	ARG	129		5.740	-2.334	28.093	1.00	15.62	
ATOM	988	NE	ARG	129		5.028		27.086		17.92	
MOTA	989	CZ	ARG	129	at No.	3.963	-3.861	27.354			1.00
MOTA	990		ARG	129	100	3.494	-3.919	28.599	1.00	15.95	
ATOM	991	NH2	ARG	129				26.382		20.39	4-13
ATOM	992	C	ARG	129		10.464	-2.327	26.468	1.00	14.38	
MOTA	993	0	ARG	129				27.097		14.82	
MOTA	994	N	ASP	130				25.478		13.77	
MOTA	995	CA	ASP	130		12.427		25.126		14.20	
MOTA		CB	ASP	130		12.055		24.111		14.89	
MOTA	997		ASP	130			-4.990	•		13.64	• • •
MOTA	998			130				23.144			
MOTA	999	002		- :130				25.058			
MOTA			ASP	130		13.548		24.561			• ••
ATOM	1001	0	ASP	130		14.166		23.554		13.46	
MOTA	1002	N	ILE	131		13.820		25.214		13.80	
MOTA	1003	CA	ILE	131		14.874	0.146	24.748		14.84	
ATOM	1004	CB	ILE	131		14.779	1.517	25.449		15.30	
MOTA	1005	CG2		131		15.976	2.412	25.055		.13.40	•
MOTA	1006	CG1		131		13.458		25.057		14.31	
MOTA	1007	CD1		131		13.093	3.399	25.917		15.14	
MOTA	1008	C	ILE	131		16.244	-0.469	25.008	1.00	14.20	. ••

FIG.11A-24

7	MOTA	1009	0	ILE	131	16.543	-0.878	26.115	1.00 14.42
1	MOTA	1010	N	LYS	132	17.054	-0.544	23.959	1.00 13.66
1	MOTA	1011	CA	LYS	132	18.405	-1.096	24.020	1.00 12.59
1	ATOM	1012	CB	LYS	132	18.376	-2.623	24.187	1.00 13.39
1	ATOM	1013	CG	LYS	132	17.494	-3.375	23.194	1.00 16.12
1	MOTA	1014	CD	LYS	132	17.518	-4.865	23.500	1.00 15.73
1	MOTA	1015	CE	LYS	132	16.670	-5.666	22.520	1.00 18.36
1	MOTA	1016	NZ	LYS	132	16.639	-7.121	22.872	1.00 16.42
4	MOTA	1017	C	LYS	132	19.084	-0.703	22.715	1.00 13.57
1	MOTA	1018	0	LYS	132	18.413	-0.351	21.749	1.00 15.48
1	ATOM	1019	N	PRO	133	20.424	-0.769	22.665	1.00 14.76
1	ATOM	1020	CD	PRO	133	21.328	-1.231	23.731	1.00 16.68
1	ATOM	1021	CA	PRO -	133	21.188		21.467	1.00 15.75
4	MOTA	1022	CB	PRO	133	22.622	-0.746	21.858	1.00 14.35
	MOTA	1023	CG	PRO	133	22.612	-0.538	23.363	1.00 16.22
	ATOM	1024	C	PRO	133	20.758	-1.055	20.162	1.00 16.05
1	MOTA	1025	0	PRO	133	20.868	-0.441	19.096	1.00 18.14
,	MOTA	1026	N	GLU	134	20.265	-2.289	20.246	1.00 15.05
	ATOM	1027	CA	GLU	134	19.820	-3.010	19.061	1.00 17.14
	ATOM	1028	CB	GLU	134	19.562	-4.488	19.404	1.00 15.98
	ATOM _	1029	CG	GLU	134	20.792	-5.246	19.898	1.00 21.80
	ATOM	1030	CD	GLU	134	20.945	-5.241	21.415	1.00 24.82
	ATOM	1031	0E1	GLU	134	20.669	-4.207	22.067	1.00 18.91
	ATOM	1032	0E2	GLU	134	21.363	-6.287	21.957	1.00 27.97
	MOTA	1033	C	GLU	134	18.554	-2.389	18.470	1.00 18.34
	ATOM	1034	0	GLU	134	18.276	-2.539	17.280	1.00 21.57
	ATOM	1035	N	ASN	135	17.785	-1.698	19.307	1.00 18.40
٠.	ATOM	1036	CA	ASN	135	16.545	-1.063	18.867	1.00 17.42
	ATOM	1037	CB	ASN	135	15.407	-1.373	19.851	1.00 16.38
	ATOM	1038	CG	ASN	135	14.881	-2.788	19.697	1.00 21.05
	ATOM	1039	OD1	ASN	135	14.895	-3.344	18.596	1.00 25.80
	ATOM	1040	ND2	ASN	135	14.397	-3.372	20.791	1.00 18.14
	ATOM	1041	C	ASN	135	16.663	0.448	18.687	1.00 17.18.
	ATOM :	1042	0	ASN	135	15.656	1.157	18.628	1.00 18.63
	ATOM	1043	N	LEU	136	17.895	0.935	18.609	1.00 15.45
	ATOM	1044	-CA	LEU	136	18.149	2.356	18.399	1.00 13.88
	ATOM	1045	CB	LEU	136	18.902	2.944	19.597	1.00 14.66
	ATOM	1046	CG	LEU	136	18.121	2.860	20.919	1.00 13:43
	ATOM	1047		LEU	136	18.9 87	3.330	22.082	1.00: 9.48
	ATOM	1048	CD2	LEU	136	16.856	3.724	20.826	
	MOTA	1049	C	LEU	136	18.984	2.416	17.122	1.00 15.09
	MOTA	1050	0	LEU	136	20.162	2.068	17.120	1.00 15.49

FIG.11A-25

MOTA	1051	N	LEU	137	18.346	2.824	16.031	1.00 16.71	
MOTA	1052		LEU	137	19.002	2.884	14.729	1.00 18.73	
MOTA	1053	CB	LEU	137	18.025	2.408	13.650	1.00 18.99	
MOTA	1054	CG	LEU	137	17.362	1.067	13.998	1.00 20.11	
MOTA	1055	CD1	LEU	137	16.443		12.863	1.00 21.15	
MOTA	1056	CD2		137	18.438		14.257	1.00 17.16	
ATOM	1057	C		137	19.532	4.274	14.400	1.00 19.64	
MOTA	1058	0	LEU	137	19.152	5.259	15.029	1.00 18.56	
MOTA	1059	N		138	20.416	4.345	13.406	1.00 20.11	,
MOTA	1060	CA		138	21.030	5.605	13.012	1.00 21.10	
MOTA	1061	CB	LEU	138	22.538			1.00 23.33	
MOTA	1062	CG	LEU	138	23.028	5.317	14.724	1.00 24.15	
MOTA	1063	CD1	LEU	138	22.444	6.368	15.650	1.00 21.36	
MOTA	1064	CD2	LEU -	138	22.620	3.921	15.169	1.00 27.70	٠.
MOTA	1065	C	LEU	138		5.872	11.526	1.00 23.67	i þ
MOTA	1066	0	LEU	138	20.963	4.959	10.707	1.00 23.81	1
MOTA	1067	N	ASP	139	20.498	7.116	11.184	1.00 24.02	
MOTA	1068	CA	ASP	139	20.298	7.481	9.784	1.00 24.89	
MOTA	1069	CB	ASP	139	19.295	8.642	9.657	1.00 23.61	
ATOM	1070	CG	ASP	139	19.861	9.974	10.120	1.00 24.18	
ATOM	1071	0D1	ASP	139	19.136	10.986	10.021	1.00 27.10	
MOTA	1072	0D2	ASP	139	21.019	10.020	10.576	1.00 24.71	
MOTA	1073	C	ASP	139	21.642	7.857	9.173	1.00 24.93	
MOTA	1074	0	ASP	139	22.687	7.630	9.781	1.00 26.58	
MOTA	1075	N	GLU	140	21.622	8.426	7.971	1.00 25.87	3
MOTA	1076	CA	GLU	140	22.857	8.808	7.296	1.00 28.66	,
MOTA	1077	CB	GLU	140	22.556	9.284	5.866	1.00 30.20	}
MOTA	1078	CG	GLU	140	21.489	10.364	more new works were a resident and a resident and a second a second and a second and a second and a second and a second an	1.00 39.13	}
ATOM	1079	CD	GLU	140	20.119	9.881	6.200		
MOTA	1080	0E1	GLU	140	19.686	8.808	5.732	1.00 50.57	7, 0
MOTA	1081	0E2	GLU	140	19.474	10.576	7.013	1.00 52.55	5 ,
MOTA	1082	C	GLU	140	23.682	9.866	8.032	1.00 28.46	5
ATOM	1083	0	GLU	140	24.905	9.914	7.882	1.00 29.94	ŀ
ATOM	1084	N	ARG	141	23.022	10.710	8.821	1.00 27.37	7.
ATOM	1085	CA	ARG	141	23.715	11.756	9.576	1.00 27.01	L
MOTA	1086	CB	ARG	141	22.942	13.076	9.479	1.00 30.09	•
ATOM	1087	CG	ARG	141	22.830	13.619	8.059	1.00 37.24	1
MOTA	1088	CD	ARG	141	22.072	14.941	7.994	1.00 44.03	3
ATOM	1089	NE	ARG	141	22.712	15.992	8.783	1.00 54.80)
ATOM	1090	CZ	ARG	141	22.445	16.242	10.052	1.00 62.47	7
ATOM	1091	NH1	ARG	- 141	21.542	15.519	10.711	1.00 60.83	L
ATOM	1092	NH2	ARG	141	23.084	17.218		•	4

FIG.11A-26

ATOM		C	ARG	141		23.891	11.362	11.045	1.00 25.81
ATOM	1094	0	ARG	141		24.141	12.206	11.909	1.00 26.51
MOTA	1095	N	ASP	142		23.779	10.066	11.312	1.00 24.59
MOTA	1096	CA	ASP	142		23.909	9.532	12.664	1.00 25.48
MOTA	1097	CB	ASP	142		25.296	9.822	13.251	1.00 25.98
MOTA	1098	CG	ASP	142		26.350	8.865	12.743	1.00 30.74
MOTA	1099	OD1	ASP	142		26.006	7.694	12.494	1.00 30.35
ATOM	1100	OD2	ASP :	142		27.521	9.272	12.608	1.00 40.27
ATOM	1101	C	ASP.	142		22.845	10.022	13.634	1.00 22.57
ATOM	1102	.0	ASP	142		23.102	10.139	14.834	1.00 22.18
ATOM	1103	N	ASN	143		21.655	10.314	13.125	1.00 22.09
MOTA	1104	CA	ASN	143		20.563	10.733	13.999	1.00 23.36
ATOM	1105	CB	ASN	143	· · ·	19.531	11.547	13.225	1.00 22.79
ATOM	1106	CG	ASN	143	٠.	20.055	12.906	12.826	1.00 26.55
MOTA	1107	OD1	ASN	143		20.119	13.240	11.644	1.00 29.98
MOTA	1108	ND2	ASN	143		20.442	13.697	13.815	1.00 24.11
MOTA	1109	Č	ASN	143		19.928	9.461	14.543	1.00 22.26
MOTA	1110	0	ASN	143		19.689	8.519	13.798	1.00 22.91
MOTA	1111	N	LEU	144		19.667	9.438	15.846	1.00 20.92
MOTA	1112	CA	LEU	144		19.084	8.268	16.494	1.00 21.74
MOTA	1113	CB	LEU	144		19.402	8.318	17.992	1.00 18.53
MOTA	1114	CG	LEU	144		18.845	7.262	18.946	1.00 20.54
MOTA	1115	CD1	LEU	144		19.807	7.095	20.113	1.00 19.77
MOTA	1116	CD2	LEU	144		17.463	7.673	19.440	1.00 21.20
MOTA	1117	C	LEU	144		17.580	8.140	16.258	1.00 20.47
MOTA	1118	0	LEU	144		16.844	9.126	,	1.00 19.22
MOTA	1119	N	LYS	145		17.140	6.909	16.000	1.00 19.53
MOTA	1120	CA	LYS	145		15.737	6.605	15.730	1.00 18.88
MOTA	1121	CB	LYS	145		15.549	6.245	14.251	1.00 24.21
MOTA	1122	CG	LYS	145		16.214	7.188	13.260	1.00 23.93
MOTA	1123	CD	LYS	145		15.328	8.369	12.951	1.00 22.67
ATOM	1124	CE	LYS	145	: :	15.970	9.275	11.913	1.00 27.57
MOTA	1125	NZ	LYS	145		15.022	10.333	11.462	1.00 27.78
MOTA	1126	C	LYS	145	٠.	15.302	5.398	16.556	1.00 15.99
MOTA	1127	0	LYS	145		15.869	4.314	16.414	1.00 16.13
MOTA	1128	N	ILE	146		14.300	5.579	17.410	1.00 16.28
MOTA	1129	CA	ILE	146		13.801	4.478	18.226	1.00 15.84
MOTA	1130	CB	ILE	146		12.849	4.993	19.319	1.00 15.02
ATOM	1131	CG	2 ILE	146		12.230	3.819	20.080	1.00 13.88
MOTA	1132	CG:	L ILE	146		13.635	5.884	20.284	1.00 17.67
MOTA	1133	CD:	1 ILE	146		1 2.781	6.595	21.324	1.00 12.64
MOTA	1134	·C	ILE	146		13.068	3.523	17.284	1.00 16.96

FIG.11A-27

MOTA	1135	0	ILE	146	12.200	3.942	16.512	1.00 17.32
MOTA	1136	N	SER -	147	13.417	2.245	17.375	1.00 16.60
MOTA	1137	CA	SER	147	12.876	1.212	16.495	1.00 17.35
MOTA	1138	CB	SER	147	14.016	0.692	15.618	1.00 16.68
MOTA	1139	OG	SER.	147	13.617	-0.411	14.821	1.00 20.69
MOTA	1140	C	SER	147	12.200	0.017	17.162	1.00 16.72
MOTA	1141	0	SER	147	12.504	-0.329	18.306	1.00 15.17
ATOM	1142	N	ASP	148	11.286	-0.602	16.414	1.00 16.05
MOTA	1143	CA	ASP	148	10.549	-1.801	16.828	1.00 18.00
ATOM	1144	CB	ASP	148	11.536	-2.919	17.200	1.00 20.31
ATOM	1145	CG	ASP	148	10.874	-4.287	17.287	1.00 25.87
MOTA	1146	OD1	ASP	148	11.601	-5.305	17.231	1.00 29.19
MOTA	1147	OD2	ASP	148	9.635	-4.349		1.00 24.90
MOTA	1148	C	ASP		137			1.00 18.24
ATOM	1149	0	ASP	148	9.887	-1.668		1.00 20.08
ATOM	1150	N	PHE	149		-1.446		
ATOM	1151		PHE	, 149	7.218	25		1.00 19.50
ATOM	1152		PHE		6.346	-0.077		
ATOM	1153		PHE	The state of the s	7.065	1.232		
ATOM	1154		PHE		7.955	1.637	The state of the s	
ATOM	1155		PHE	149	6.932	2.014		1.00 15.99
ATOM	1156	4.1		149	8.712	2.805	The state of the s	1.00 19.45
MOTA		CE2		149		3.184		1.00 20.50
ATOM		CZ			8.576	and the second s	18.568	
MOTA	1159		PHE	149			18.780	1.00 20.77
MOTA	1160		PHE	149	5.235		19.187	A. A.
ATOM	1161		GLY	150			18.539	
MOTA	1162		GLY		6.361	-4.942	18.725	1.00 21.69
MOTA	1163				6.002	and the second second	20.176	
MOTA	1164		GLY	• • • • • • • • • • • • • • • • • • • •	5.111		20.449	1.00 24.57
MOTA				151	6.687		21.111	1.00 19.98
MOTA				151				1.00 21.36
MOTA				151			•	1.00 18.98
MOTA								1.00 24.33
MOTA		CD1						1.00 25.19
MOTA		CD2					23.440	
MOTA	1171		LEU		·		23.195	
MOTA	1172		LEU		5.517		24.389	,
MOTA	1173		ALA		5.640		22.413	
MOTA	1174		ALA		5.102		22.915	
MOTA		CB	ALA					1.00 16.88
MOTA	1176	C	ALA	152	3.627	-1.290	23.285	1.00 19.46

FIG.11A-28

 ·					•				•		
 ATOM	1177	0	ALA	152	2.8	395 -2	. 129	22.758	1.00	21.66	•
MOTA	1178	N	THR	153	3.3	192 -0	.418	24.189	1.00	19.05	
MOTA	1179	CA	THR	153	1.3	796 -0	. 397	24.593	1.00	18.72	
MOTA	1180	CB	THR	153	1.5	509 -1	.442	25.712	1.00	18.18	
MOTA	1181	OG1	THR	153	0.0	090 -1	.652	25.809	1.00	16.70	
MOTA	1182	CG2	THR	153	2.	038 -0	. 970	27.071	1.00	16.94	
ATOM	1183	C	THR	153	1.3	396 1	.000	25.056	1.00	19.47	
MOTA	1184	0	THR	153	2.	244 1	.853	25.325	1.00	17.58	
MOTA	1185	N	VAL	154	G.	096 1	. 249	25.112	1.00	21.41	
MOTA	1186	CA	VAL	154	-0.	401 2	.543	25.547	1.00	22.13	
MOTA	1187	CB	VAL :	154	-1.	765 2	.863	24.877	1.00	26.46	
ATOM	1188	CG1	VAL	154	-2.	295 4	.213	25.354	1.00	28.65	
MOTA	1189	CG2	VAL :	154.	:-1.	600 2	.873	23.367	1.00	24.98	
MOTA	1190	C	VAL	154	-0.	559 2	.472	27.056	1.00	21.16	
ATOM	1191	0	VAL	154	-1.	195 1	.553	27.577	1.00	21.97	
MOTA	1192	N	PHE	155	0.	047 3	.416	27.770	1.00	19.24	
MOTA	1193	CA	PHE	155	-0.	061 3	.414	29.220	1.00	18.72	
MOTA	1194	CB	PHE	155	1.	322 3	.426	29.889	1.00	19.67	
MOTA	1195	CG	PHE	155	2.	055 4	.721	29.748	1.00	17.34	
MOTA	1196	CD1	PHE	155	2.	843 4	.972	28.628	1.00	13.52	
MOTA	1197	CD2	PHE	155	1.	924 5	.711	30.716	1.00	16.84	
MOTA	1198	CE1	PHE	155			. 191	28.470	1.00	13.59	
MOTA	1199	CE2	PHE	155	2.	565 6	.944	30.570	1.00	16.30	
MOTA	1200	CZ	PHE	- 155	3.	350 7	1.187	29.445	1.00	15.78	
MOTA	1201	C	PHE	155	-0.	889 4	.590	29.717	1.00	20.17	
MOTA	1202	0	PHE	155	-1.	170 4	.696	30.907	1.00	20.34	
ATOM	1203	N	ARG	156	-1.	259 5	489	28.812	1.00	19.17	
ATOM	1204	CA	ARG	156	-2.	096 6	.622	29.204	1.00	18.98	
ATOM	1205	CB	ARG	156	-1.	282 7	.904	29.388	1.00	17.96	
ATOM	1206	CG		156	-2.		.008	30.101	1.00	20.91	
ATOM	1207	CD	ARG	156	-1.	432 10	.382	29.971	1.00	26.19	
ATOM	1208	NE	ARG	156	-0.	049 10	.410	30.438	1.00	25.61	
MOTA	1209		ARG	156	1.	002 10	.600	29.642	1.00	20.60	
ATOM	1210	NH1	ARG	156	0.	830 10	.774	28.340	1.00	19.60	
MOTA	1211	NH2	ARG	156	2.	226 10	.628	30.150	1.00	18.11	
MOTA	1212	C	ARG	156	· -3.	134 6	.847 °	28.122	1.00	21.82	
ATOM	1213	0	ARG	156	-2.	802 7	.039	26.954	1.00	21.83	
ATOM	1214	N	TYR	157	-4.	398 6	.824	28.521	1.00	21.32	
ATOM	1215	CA	TYR	157	-5.	493 7	.016	27.584	1.00	19.71	
MOTA	1216	CB	TYR	157	-6.	101 5	663	27.218	1.00	18.64	
MOTA	1217	CG	TYR	157	-6.	960 5	5.702	25.983		23.71	
MOTA	1218	CD1	TYR	157	-6.	384 5	.726	24.712		21.76	

FIG.11A-29

MOTA	1219	CE1 1	ΓYR	157		-7.174	5.767	23.566	1.00 2	25.10	
MOTA	1220	CD2 T	ΓYR	157		-8.350	5.719	26.081	1.00	19.65	
MOTA	1221	CE2	ΓΥR	157	•	-9.147	5.756	24.946	1.00	19.21	•
MOTA	1222	CZ	ΓYR	157		-8.559	5.780	23.693	1.00	22.17	
MOTA	1223	OH 7	TYR	157	· •	-9.347	5.818	22.566	1.00	25.91	
MOTA	1224	C	TYR	157		-6.533	7.882	28.282	1.00	18.10	
ATOM	1225	0	TYR	157	•	-6.851	7.651	29.449	1.00	18.79	:
ATOM			ASN .	158		-7.045	8.881	27.571	1.00	19.43	
MOTA	1227	CA.	ASN	158	,	-8.041	9.797	28.130	1.00	22.45	
MOTA	1228	CB	ASN	158		-9.375	9.068	28.342	1.00	17.95	
MOTA	1229	CG	ASN	158		10.134	8.861	27.046	1.00	15.00	
MOTA	1230	001	ASN	158		-11.036	8.025	26.968	1.00	21.50	. •
ATOM	1231	ND2	ASN -	: 158		-9.777	9.620	26.018		14.33	
ATOM	1232	C	ASN	158		-7.565	10.417	29.442	1.00	24.59	•
ATOM	1233	0	ASN	158		-8.339	10.591	30.383	1.00	25.26	
ATOM	1234	N	ASN	159		-6.272	10.731	29.482	1.00	27.27	
ATOM	1235	CA	ASN	159	Š.	-5.624	11.353	30.630		28.47	
ATOM	1236	CB	ASN	159		-6.285	12.702	30.934		34.81	
ATOM	1237	CG	ASN	159		-5.380	13.624	31.730		46.84	. 0
ATOM	1238	OD1	ASN	159	e.	-4.243	13.884	31.332		48.82	
ATOM	1239	ND2	ASN	159		-5.880	14.126			48.82	
ATOM	1240	C	ASN	159		-5.597		31.889		27.91	
ATOM	1241	0	ASN	159		-5.381	10.991	32.993		30.07	
MOTA	1242	N	ARG	160		-5.818		31.725	and the second of the second	24.35	
ATOM	1243	CA	ARG	160	7. C	-5.788	•	32.854		23.07	
ATOM	1244	CB	ARG	160		-7.104		32.961		24.32	
MOTA	1245	CG	ARG	160	: ⁻	-8.050		34.040		31.06	٠.,
MOTA	1246	CD	ARG	160		-7.472		35.429		36.21	
ATOM	1247	NE	ARG	160		-8.462		36.479		49.83	
ATOM	1248		ARG	160		-8.983	9.174			58.03	
MOTA	1249					-8.608		36.135		61.87	٠
ATOM				160	•	-9.887		37.758		59.82	
MOTA	1251	C		160		-4.639	•	32.648		22.67	:
ATOM	1252	^ O	ARG	160				31.618		20.87	٠
ATOM	1253	N	GLU	161	•			33.630		22.22	
ATOM	1254	CA	GLU	161				33.527		22.53	
ATOM	1255	CB	GLU	161				34.429	•	22.92	
MOTA	1256		GLU	161			•	34.340		21.95	
MOTA	1257		GLU	161			•	35.305		24.35	
ATOM	1258		GLU	161	٠		5.637			20.96	
ATOM	1259	OE2	GLU	. 161				36.095		25.52	
MOTA	1260	C	GLU	161		-3.007	4.885	33.926	1.00	21.71	

FIG.11A-30

ATOM	1261	0	GLU	161	-3.755	4.683	34.885	1.00 23.04
ATOM	1262	N	ARG	162	-2.503	3.909	33.182	1.00 21.16
MOTA	1263	CA	ARG	162	-2.754	2.505	33.465	1.00 21.51
MOTA	1264	CB	ARG	162	-3.207	1.781	32.191	1.00 26.46
MOTA	1265	CG	ARG	162	-3.326	0.274	32.326	1.00 33.90
ATOM	1266	CD	ARG	162	-3.916	-0.347	31.061	1.00 44.41
MOTA	1267	NE	ARG	162	-3.035	-0.230	29.898	1.00 54.96
MOTA	1268	CZ	ARG	162	-2.050	-1.077	29.612	1.00 52.17
ATOM	1269	NH1	ARG	162	-1.303	-0.884	28.534	1.00 48.31
MOTA	1270	NH2	ARG	162	-1.816	-2.123	30.392	1.00 49.02
MOTA	1271	C	ARG	162	-1.442	1.892	33.957	1.00 21.44
ATOM	1272	0	ARG	162	-0.405	2.058	33.319	1.00 20.36
ATOM	1273	N	LEU.	163	-1.481	1.215	35.098	1.00 20.22
MOTA	1274	CA	LEU	163	-0.279	0.573	35.623	1.00 21.99
ATOM	1275	CB	LEU	163	-0.448	0.226	37.100	1.00 22.03
_ATOM	1276	CG	LEU	163	-0.661	1.398	38.057	1.00 23.54
ATOM	1277	CD1	LEU	163	-1.002	0.862	39.439	1.00 21.82
ATOM	1278	CD2	LEU	163	0.598	2.269	38.100	1.00 23.24
ATOM	1279	C	LEU	163	-0.051	-0.699	34.823	1.00 22.61
ATOM	1280	0	LEU	163	-1.000	-1.362	34.411	1.00 23.66
MOTA	1281	N .	LEU	164	1.211	-1.045	34.604	1.00 21.45
MOTA	1282	CA	LEU	164	1.526	-2.245	33.839	1.00 19.50
ATOM	1283	CB	LEU	164	2.699	-1.966	32.898	1.00 19.82
MOTA	1284	CG	LEU	164	2.524	-0.748	31.991	1.00 21.21
ATOM	1285	CD1	LEU	164	3.741	-0.606	31.096	1.00 23.59
ATOM	1286	CD2		164	1.260	-0.897	31.166	1.00 21.35
ATOM	1287	C.	LEU	164	1.887	-3.402	34.752	1.00 17.33
ATOM	1288	0	LEU	164	2.254	-3.194	35.909	1.00 16.78
ATOM	1289	N	ASN	165	1.784	-4.621	34.222	1.00 17.32
ATOM	1290	CA	ASN	165	2.139	-5.818	34.978	1.00 19.46
MOTA	1291	CB	ASN	165	0.898	-6.443	35.622	1.00 22.03
ATOM	1292		ASN	165	-0.189	-6.740	34.611	1.00 24.96
MOTA	1293		ASN	165	-1.219	-6.065	34.574	1.00 31.70
ATOM	1294	ND2	ASN	165	0.037	-7.748	33.776	1.00 22.31
ATOM	1295	. C	ASN	165	2.816	-6.855	34.084	1.00 19.59
ATOM	1296	0	ASN	165	3.349	-7.849	34.569	1.00 21.55
ATOM	1297	N	LYS	166	2.804	-6.625	32.778	1.00 20.65
MOTA	1298	CA	LYS	166	3.425 .	-7.570	31.854	1.00 23.25
ATOM	1299	CB	LYS	166	3.029	-7.232	30.414	1.00 25.58
ATOM	1300	CG	LYS	166	3.605	-8.164	29.356	1.00 28.68
ATOM	1301	CD	LYS	166	3.109	-7.776		1.00 34.56
ATOM	1302	CE	LYS	166	3.602	-8.742		1.00 40.12

FIG.11A-31

ATOV	1202	ALŻ	ive	166		F 000	0.750	ÖC 011	1 00 47 02
ATON	1303	NZ	LYS	166	•	5.089	-8.750	26.811	1.00 47.83
MOTA	1304	C	LYS	166	•	4.949	-7.569	31.982	1.00 22.75
MOTA	1305	0	LYS	166		5.594	-6.523	31.884	1.00 20.68
MOTA	1306		MET	167	. 0	5.523	-8.741	32.230	1.00 22.74
ATOM	1307	CA	MET	167		6.973	-8.835	32.320	1.00 23.09
MOTA	1308	CB	MET	167	42.5		-10.040	33.163	
MOTA	1309	CG	MET	167		7.362			1.00 24.94
ATOM	1310	SD		167				35.618	700
MOTA	1311		MET .	167			-12.283		1.00 40.11
MOTA	1312		MET	167			-8.985	the state of the s	
ATOM	1313	0	MET	167				·	1.00 24.88
ATOM	1314		CYS	168			-7.989		
ATOM	1315	CA		168	10		-8.018		1.00 19.64
ATOM	1316	CB	CYS					28.061	
MOTA	1317		CYS	168			-5.932		1.00 25.33
ATOM	1318	C	CYS	168		7			1.00 19.48
MOTA	1319	0	CYS	168			-6.341		
MOTA	1320	N	GLY	169					1.00 18.34
ATOM	1321	CA	GLY	169		11.867	-6.422		1.00 16.84
MOTA	1322	C	GLY	169		13.056	-7.347	27.473	
MOTA	1323	0	GLY	169		12.910	-8.446	26.932	1.00 17.29
MOTA	1324	N	THR	170		14.225	-6.898	27.922	1.00 16.83
MOTA	1325	CA	THR	170		15.473	-7.649	27.811	1.00 17.19
MOTA	1326	CB	THR	170		16.343	-7.057	26.678	1.00 16.59
MOTA	1327	OG:	L THR	170		15.593	-7.087	25.453	1.00 16.29
MOTA	1328	CG	2 THR	170		17.606	-7.871	26.483	1.00 17.54
MOTA	1329	C	THR	170	i Den e	16.160	-7.520	29.176	1.00 15.39
MOTA	1330	0	THR	170		16.494	-6.416	29.608	and the second s
MOTA	1331	N	LEU	171	1 y	16.374	-8.658	29.838	1.00 15.78
MOTA	1332	CA	LEU	171		16.938	-8.697	31.190	1.00 15.62
MOTA	1333	CB	LEU	171					1.00 17.62
MOTA			LEU	171		16.781	-10.963	32.621	1.00 24.25
MOTA	1335	CD	1 LEU	171	: •	15.469	-10.373	33.131	1.00 20.87
MOTA	1336	CD	2 LEU	171		16.577	-12.390	32.116	1.00 14.78
MOTA	1337	C	LEU	· 171		18.007	-7.675	31.615	1.00 15.63
MOTA	1338		LEU	171		17.835	-6.989	32.625	1.00 14.39
MOTA	1339		PRO	172		19.123	-7.559	30.872	1.00 15.88
ATOM	1340		• *	172		19.564	-8.355	29.713	1.00 18.04
MOTA	1341	CA		172		20.156			
MOTA	1342			172		21.268	•		
MOTA	1343			172		21.060			
MOTA	1344		PRO	172		19.689			
		_							·

FIG.11A-32

ATOM	1045	, , , , , , , , , , , , , , , , , , ,	170		00.055			
MOTA	1345	O PRO	172		20.268	-4.291	31.972	1.00 18.87
MOTA	1346	N TYR	•		18.630	-4.852	30.532	1.00 15.37
ATOM	1347	CA TYR	173		18.073	-3.506	30.421	1.00 14.75
MOTA	1348	CB TYR	173		17.757	-3.218	28.950	1.00 13.39
MOTA	1349	CG TYR	173		18.954	-3.298	28:046	1.00 14.82
MOTA	1350	CD1 TYR	173		19.745	-2.182	27.811	1.00 15.47
MOTA	1351	CE1 TYR	173		20.872	-2.255	26.993	1.00 20.45
MOTA	1352	CD2 TYR	173		19.314	-4.503	27.438	1.00 19.99
MOTA	1353	CE2 TYR	173		20.435	-4.585	26.617	1.00 23.28
MOTA	1354	CZ TYR	173		21.208	-3.455	26.401	1.00 20.15
MOTA	1355	OH TYR	173		22.317	-3.523	25.586	1.00 23.35
MOTA	1356	C TYR	173		16.795	-3.271	31.223	1.00 14.06
ATOM	1357	O TYR	· 173		16.336	-2.135	31.351	1.00 13.16
MOTA	1358	N VAL	174		16.212	-4.328	31.771	1.00 15.36
MOTA	1359	CA VAL	174		14.950	-4.171	32.485	1.00 15.69
MOTA	1360	CB VAL	174		14.183	-5.529	32.498	1.00 18.37
MOTA	1361	CG1 VAL	174		14.686	-6.421	33.634	1.00 16.95
MOTA	1362	CG2 VAL	174		12.689	-5.284	32.590	1.00 20.81
MOTA	1363	C VAL	174		15.083	-3.596	33.909	1.00 14.76
MOTA	1364	O YAL	174		16.048	-3.875	34.616	1.00 14.52
MOTA	1365	N ALA	175		14.109	-2.778	34.302	1.00 14.40
ATOM	1366	CA ALA	175		14.099	-2.152	35.628	1.00 14.61
MOTA	1367	CB ALA	175		13.044	-1.055	35.669	1.00 15.96
MOTA	1368	C ALA	- 175		13.830	-3.185	36.729	1.00 14.55
MOTA	1369	O ALA	175		13.079	-4.130		1.00 14.73
MOTA	1370	N PRO	176		14.435	-3.001	37.912	
MOTA	1371	CD PRO	176		15.321	-1.891	38.303	1.00 16.61
ATOM	1372	CA PRO	176		14.247	-3.941	39.022	1.00 15.95
MOTA	1373	CB PRO	176		15.154	-3.372	40.120	1.00 18.53
MOTA	1374	CG PRO	176		15.200	-1.896	39.812	1.00 17.80
ATOM	1375	C PRO	176	::	12.805	-4.157	39.487	
ATOM	1376	O PRO	176		12.456		39.923	
ATOM	1377	N GLU	177		11.958			1.00 17.70
ATOM	1378	CA GLU	177			-3.294		1.00 19.33
MOTA	1379		177		9.831	-1.954		1.00 19.43
ATOM	1380	CG GLU	177		9.711		38.479	
MOTA	1381	CD GLU	177		10.866	-0.291		1.00 18.08
ATOM	1382	OE1 GLU	177		10.672		37.389	1.00 15.25
ATOM	1383	OE2 GLU			11.962		38.775	1.00 19.51
ATOM	1384	C GLU			9.815		38.977	
ATOM	1385	0 GLU	177		8.877			1.00 19.97
ATOM	1386	N LEU	178		10.214	-4.485		1.00 18.60
, O	1000		1,0		70.574	7.705	J1.121	7.00 TO.00

FIG.11A-33

						Sec. 1				
MOTA	1387	CA I	LEU.	178		9.540	-5.448	36.861	1.00 21.80)
MOTA		CB I	LEU	178		10.037	-5.283	35.412	1.00 26.36	;
MOTA			LEU	178		9.551	-6.196	34.281	1.00 30.81	•
MOTA	1390	CD1	LEU	178		10.271	-7.531	34.349	1.00 32.00)
MOTA		CD2	LEU	178	-	8.053	-6.389	34.371	1.00 34.42	2
ATOM			LEU	178		9.789	-6.866	37.379	1.00 21.49	
ATOM			LEU	178	•	8.987	-7.776	37.148	1.00 22.54	1
ATOM	•		LEU	179	• •	10.886	-7.051	38.107	1.00 22.22	2 .
MOTA			LEU	179		11.213	-8.365	38.648	1.00 23.39	9
ATOM		CB	LEU	179	٠.	12.719	-8.621	38.558	1.00 23.03	3
MOTA		CG	LEU	179		13.416	-8.495	37.200	1.00 26.29	9
MOTA	1398	CD1	LEU.	179	**	14.903	-8.733	37.390	1.00 24.6	7
MOTA	1399	_		179	,	12.837	-9.491	36.204	1.00 26.5	7
MOTA	1400	C	LEU	179		10.770	-8.558	40.096	1.00 24.8	4
MOTA	1401	0	LEU	179		10.847	-9.667	40.627	1.00 26.9	2
MOTA	1402	N	LYS	180	, in	10.295	-7.504	40.746	1.00 23.3	5
MOTA	1403	CA	LYS	180		9.908	-7.666	42.143	1.00 25.6	9
ATOM	1404	CB	LYS	180	j.	10.916	-6.949	43.044	1.00 31.3	9
ATOM	1405	CG	LYS	180		11.002	-5.452	42.823	1.00 40.6	1
MOTA	1406	CD	LYS	180	4	12.048	-4.816	43.737	1.00 49.3	8
MOTA	1407	CE	LYS	180	4	13.441	-5.362	43.457	1.00 56.3	
MOTA	1408	NZ	LYS	180		14.482	-4.726		1.00 63.7	
MOTA	1409	C	LYS	180	. 7	8.508	-7.228	**	1.00 26.8	
MOTA	1410	0	LYS	180		8.025	-7.586		1.00 27.5	
MOTA	1411	N -	ARG	181	ý.º	7.849	-6.471		1.00 25.9	
MOTA	1412	CA	ARG	181	.:	6.507		41.953	1.00 23.6	
MOTA	1413	CB	ARG	181		6.515		42.013		
MOTA	1414	_CG_	ARG			7.886	-3.864		1.00 23.4	
MOTA	1415	CD	ARG	181		7.952	-3.096		1.00 28.2	
MOTA	1416		ARG	181		7.769	-3.932	44.835	1.00 26.3	
ATOM	1417			- 181		8.303		46.032	1.00 25.0	
ATOM	1418			181		8.059			1.00 21.2	
MOTA	1419	NH2	ARG			9.096		46.221		
MOTA	1420	C	ARG	181		5.489		40.921		
ATOM	1421	0	ARG	181		5.813		39.743		
MOTA	1422	* N	ARG	182		4.257		41.362		
MOTA	1423	CA	ARG	182	•	3.214	-7.141	40.452		
ATOM	1424	CB.	ARG	182	•	2	-7.550	41.229		
ATOM	1425	CG	ARG		•			40.382		
ATOM	1426	CD	ARG		•.		-8.462			
MOTA	1427	NE	ARG	182		-1.032		41.265	•	
ATOM	1428	CZ	ARG	182		-2.245	-6.998	41.781	1.00 66.4	47

FIG.11A-34

ATOM	1429	NH1 A			-2.954	-8.049	42.170	1.00 73.47
ATOM	1430	NH2 A	•		-2.750	-5.778	41.905	1.00 66.10
ATOM	1431		NRG 18		2.852	-6.046	39.450	1.00 25.07
MOTA	1432		ARG 18		2.667	-6.320	38.261	1.00 25.61
MOTA	1433		LU 18		2.744	-4.812	39.936	1.00 23.08
MOTA	1434		GLU 18		2.406	-3.673	39.085	1.00 23.45
MOTA	1435		alu 18	3	1.067	-3.059	39.501	1.00 21.25
MOTA	1436	CG 6	GLU 18	3	-0.147	-3.899	39.187	1.00 24.48
ATOM	1437		alu 18	3	-1.423	-3.181	39.569	1.00 30.66
MOTA	1438	0E1 6	alu 18	13	-1.611	-2.902	40.771	1.00 34.79
MOTA	1439	0E2 6	SLU 18	13	-2.228	-2.883	38.666	1.00 30.34
MOTA	1440	C 6	SLU 18	3	3.482	-2.600	39.169	1.00 21.06
ATOM ·	1441	···0 6	SLU 18	33	4.209	-2.512	40.158	1.00 21.34
MOTA	1442	N F	PHE 18	4	3.565	-1.768	38.137	1.00 18.15
MOTA	1443	CA F	PHE 18	34	4.567	-0.717	38.105	1.00 15.58
MOTA	1444	CB F	PHE 18	34	5.945	-1.346	37.819	1.00 16.74
MOTA	1445	CG F	PHE 18	34	5.926	-2.381	36.726	1.00 14.62
MOTA	1446	CD1 I	PHE 18	34	5.951	-2.005	35.392	1.00 18.17
MOTA	1447	CD2 I	PHE 18	34.	5.815	-3.739	37.036	1.00 17.28
MOTA	1448	CE1	PHE 18	34	5.860	-2.959	34.375	1.00 20.20
MOTA	1449	CE2	PHE 18	34	5.721	-4.698	36.029	1.00 16.96
ATOM	1450	CZ I	PHE 18	34	5.741	-4.306	34.696	1.00 17.04
MOTA	1451	C	PHE 18	34	4.222	0.353	37.067	1.00 16.19
MOTA	1452	0 0 I	PHE 18	34	3.506	0.084	36.096	1.00 15.45
MOTA	1453	N I	HIS 18	35	4.707	1.569	37.298	1.00 16.14
ATOM	1454	CA I	HIS 18	35	4.499	2.688	36.380	1.00 17.03
MOTA	1455		HIS 18	35	4.911	3.998	37.057	1.00 15.20
MOTA	1456	CG	HIS 18	35	3.954	4.462	38.110	1.00 17.47
MOTA	1457	CD2		35	4.016	4.403	39.463	1.00 16.97
MOTA	1458	ND1		35	2.755	5.074	37.808	1.00 17.76
MOTA	1459	CE1		35	2.122	5.373	38.930	1.00 15.00
MOTA	1460			35	2.866	4.978	39.948	1.00 16.42
MOTA	1461			35	5.346	2.468	35.121	1.00 16.88
MOTA	1462			35 🐇 .	6.489	2.023	35.202	1.00 15.15
MOTA	1463	N A	ALA 18	36	4.789	2.789	33.959	1.00 15.23
MOTA	1464			36	5.500	2.584	32.696	1.00 14.49
MOTA	1465	CB A	ALA 18	36	4.543	2.773	31.529	1.00 11.65
MOTA	1466			36 ·	6.719	3.472	32.469	1.00 15.84
MOTA	1467			36	7.768	2.999	32.039	1.00 13.75
MOTA	1468			37	6.579	4.760	32.747	1.00 13.01
MOTA	1469			37	7.665	5.694	32.475	1.00 14.78
MOTA	1470	CB (GLU 18	37	7.190	7.118	32.758	1.00 13.83

FIG.11A-35

			40.4	•				
MOTA	1471	CG	GLU	187	6.131	7.564	31.755	1.00 14.84
MOTA	1472	CD	GLU	187	5.476	8.860	32.155	1.00 17.06
MOTA	1473	OE1	GLU	187	5.783	9.898	31.537	1.00 17.90
MOTA	1474	0E2	GLU	187	4.669	8.836	33.101	1.00 25.47
MOTA	1475	С	GLU	187	9.023	5.420	33.119	1.00 13.36
MOTA	1476	0	GLU	1 87	10.044	5.468	32.435	1.00 14.03
MOTA	1477	N	PRO	188	9.064	5.134	34.427	1.00 12.65
MOTA	1478	CD.	PRO	188	8.004	5.222	35.448	1.00 12.15
MOTA	1479	CA	PRO	188	10.369	4.868	35.042	1.00 11.62
MOTA	1480	CB	PRO	188	10.029	4.690	36.532	1.00 13.87
MOTA	1481	CG	PRO -	188	8.799	5.543	36.707	1.00 12.10
ATOM	1482	C	PRO	188	11.079	3.639	34.471	1.00 11.69
MOTA	1483	0	PRO	188	12.302	3.525	34.575	1.00 13.08
ATOM	1484	N	VAL	189	10.324	2.709	33.878	1.00 12.14
MOTA	1485	CA	VAL	189	10.934	1.508	33.313	1.00 12.36
MOTA	1486	CB	VAL	189	9.845	0.440	32.946	1.00 11.17
MOTA	1487	CG1	VAL	189	10.485	-0.758		1.00 11.29
MOTA	1488	CG2	VAL	189	9.135		34.207	1.00 12.20
MOTA	1489	C	VAL	189	11.746	1.907	32.069	1.00 14.29
MOTA	1490	0	VAL	189	12.877	1.442		1.00 13.97
MOTA	1491	N	ASP	190	11.180	2.781		1.00 13.89
MOTA	1492	CA	ASP	190	11.882	3.237		1.00 13.47
MOTA	1493	CB	ASP	190	10.952	•		1.00 15.20
MOTA	1494		ASP	190	10.078	3.154		1.00 17.71
MOTA	1495	OD 1		190	10.434	1.981		1.00 16.91
MOTA	1496	OD2	ASP	1 9 0	9.037	3.652		1.00 17.19
MOTA	1497	C	ASP	19 0	13.062	4.124		1.00 13.29
MOTA	1498	0_	_ASP_	190	14.109	4.135		1.00 11.95
MOTA	1499	N	VAL	191	12.903	4.870	•	
MOTA	1500		VAL	191	14.009	5.716	31.988	1.00 14.45
MOTA	1501		VAL	191	13.602	6.603		
MOTA	1502		L VAL	191	14.842	7.202		
MOTA	1503	CG	2 VAL		12.688	7.727		
MOTA	1504	C		191	15.203	4.840		1.00 13.08
MOTA	1505	0	VAL	191	16.346			,
MOTA	1506	N	TRP	192	14.921	3.756		
MOTA	1507	CA	TRP		15.958	2.833		1.00 12.03
MOTA	1508	CB	TRP		15.322	1.727		1.00 9.46
MOTA	1509		TRP		16.294	0.677		1.00 12.75
MOTA	1510		2 TRP	i contract of the contract of	16.899	0.563		
MOTA	1511		2 TRP		17.767	-0.550		
MOTA	1512	CE	3 TRP	192	16.789	1.294	37.338	1.00 12.63

FIG.11A-36

1513	CD1	TRP	192		16.804	-0.342	34.098	1.00 12.01
1514	NE1	TRP	192		17.691	-1.086		1.00 12.49
1515	CZ2	TRP	192		18.525	-0.952	37.215	1.00 12.51
1516	CZ3	TRP	192		17.537	0.894	38.439	1.00 14.40
1517	CH2	TRP	192		18.396	-0.221	38.368	1.00 12.56
1518	C .	TRP	192		16.713	2.226		1.00 12.79
1519	0	TRP	192		17.947	2.240	32.345	1.00 12.82
1520	N :	SER	193		15.991	1.706		1.00 13.03
1521	CA :	SER	193		16.676	1.118		
1522	CB	SER	193		15.672	0.467	29.263	1.00 11.41
1523	OG	SER	193		14.658	1.368	28.864	1.00 14.36
1524	C	SER	193		17.523	2.175	29.506	1.00 13.32
1525	0	SER	193		18.582	1.866	28.973	1.00 12.69
1526	N	CYS	194		17.064	3.420	29,471	1.00 12.77
1527	CA	CYS	194		17.886	4.463	28.840	1.00 13.01
1528	CB	CYS	194	 •	17.136	5.799	28.793	1.00 11.84
1529	SG	CYS	194		15.813	5.829	27.558	1.00 12.71
1530	C	CYS	194		19.195	4.643	29.624	1.00 11.67
1531	0	CYS	194		20.223	4.970	29.050	1.00 13.09
1532	N	GLY	195		19.137	4.424	30.934	1.00 11.41
1533	CA	GLY	195		20.324	4.541	31.776	1.00 12.03
1534	С	GLY	195		21.311	3.421	31.480	1.00 12.90
1535	0	GLY	195		22.529	3.624	31.491	1.00 12.32
1536	: - N	ILE	196		20.792	2.225	31.223	1.00 13.85
1537	CA	ILE.	196		21.673	1.100	30.899	1.00 15.56
1538	CB	ILE	196		20.896	-0.240	30.942	1.00 21.20
1539	CG2	ILE	196 ·		19.649	-0.143	30.132	1.00 21.27
1540	CG1	ILE	196		21.763	-1.380	30.415	1.00 20.74
1541	CD1	ILE	196	-	22.970	-1.620	31.237	1.00 36.22
1542	. C	ILE	196		22.294	1.345	29.516	1.00 13.15
1543	· 0·	ILE	196		23.459	1.009	29.277	1.00 12.36
1544	· N	VAL	197	. -	21.527	1.941	28.603	1.00 12.90
1545	CA	VAL	197		22.054	2.257	27.278	1.00 13.36
1546	CB	VAL	197		20.957	2.852	26.349	1.00 13.82
1547	CG1	VAL	197		21.593	3.495	25.106	1.00 12.22
1548	CG2	VAL	197		19.986	1.740	25.929	1.00 13.53
1549	C	VAL	197		23.193	3.270	27.438	1.00 14.88
1550	0	VAL	197		24.220	3.168	26.767	1.00 16.68
1551	· N	LEU.	· 198		23.026	4.231	28.344	
1552	CA	LEU	· 198		24.060	5.244	28.561	
1553	CB	LEU	: 198		23.579	6.306	29.552	
1554	CG	LEU	198		23.930	7.793	29.353	1.00 21.66
	1514 1515 1516 1517 1518 1519 1520 1521 1522 1523 1524 1525 1526 1527 1528 1529 1530 1531 1532 1533 1534 1535 1536 1537 1538 1539 1540 1541 1542 1543 1544 1545 1546 1547 1548 1549 1550 1551 1552 1553	1514 NE1 1515 CZ2 1516 CZ3 1517 CH2 1518 C 1519 O 1520 N 1521 CA 1522 CB 1523 OG 1524 C 1525 O 1526 N 1527 CA 1528 CB 1529 SG 1530 C 1531 O 1532 N 1532 N 1533 CA 1534 C 1535 O 1536 N 1537 CA 1538 CB 1539 CG2 1540 CG1 1541 CD1 1542 C 1543 O 1544 N 1545 CA 1546 CB 1547 CG1 1548 CG2 1549 C 1550 O 1551 N 1552 CA 1553 CB	1514 NE1 TRP 1515 CZ2 TRP 1516 CZ3 TRP 1517 CH2 TRP 1518 C TRP 1519 O TRP 1520 N SER 1521 CA SER 1522 CB SER 1523 OG SER 1524 C SER 1525 O SER 1526 N CYS 1527 CA CYS 1528 CB CYS 1529 SG CYS 1529 SG CYS 1530 C CYS 1531 O CYS 1532 N GLY 1533 CA GLY 1533 CA GLY 1534 C GLY 1535 O GLY 1535 O GLY 1536 N ILE 1537 CA ILE 1537 CA ILE 1538 CB ILE 1539 CG2 ILE 1540 CG1 ILE 1541 CD1 ILE 1542 C ILE 1543 O ILE 1544 N VAL 1545 CA VAL 1546 CB VAL 1546 CB VAL 1546 CB VAL 1547 CG1 VAL 1548 CG2 VAL 1549 C VAL 1549 C VAL 1550 O VAL 1551 N LEU 1552 CA LEU 1552 CA LEU	1514 NE1 TRP 192 1515 CZ2 TRP 192 1516 CZ3 TRP 192 1517 CH2 TRP 192 1518 C TRP 192 1519 O TRP 192 1520 N SER 193 1521 CA SER 193 1522 CB SER 193 1523 OG SER 193 1524 C SER 193 1525 O SER 193 1526 N CYS 194 1527 CA CYS 194 1529 SG CYS 194 1529 SG CYS 194 1530 C CYS 194 1531 O CYS 194 1532 N GLY 195 1533 CA GLY 195 1534 C GLY 195 1535 O GLY 195 1536 N ILE 196 1537 CA ILE 196 1537 CA ILE 196 1539 CG2 ILE 196 1539 CG2 ILE 196 1540 CG1 ILE 196 1541 CD1 ILE 196 1542 C ILE 196 1543 O ILE 196 1544 N VAL 197 1545 CA VAL 197 1546 CB VAL 197 1548 CG2 VAL 197 1549 C VAL 197 1549 C VAL 197 1550 O VAL 197 1551 N LEU 198 1552 CA LEU 198 1552 CA LEU 198	1514 NE1 TRP 192 1515 CZ2 TRP 192 1516 CZ3 TRP 192 1517 CH2 TRP 192 1518 C TRP 192 1519 O TRP 192 1520 N SER 193 1521 CA SER 193 1522 CB SER 193 1523 OG SER 193 1524 C SER 193 1525 O SER 193 1526 N CYS 194 1527 CA CYS 194 1529 SG CYS 194 1529 SG CYS 194 1530 C CYS 194 1531 O CYS 194 1532 N GLY 195 1533 CA GLY 195 1534 C GLY 195 1535 O GLY 195 1536 N ILE 196 1537 CA ILE 196 1537 CA ILE 196 1539 CG2 ILE 196 1540 CG1 ILE 196 1540 CG1 ILE 196 1541 CD1 ILE 196 1542 C ILE 196 1543 O ILE 196 1544 N VAL 197 1545 CA VAL 197 1546 CB VAL 197 1546 CB VAL 197 1547 CG1 VAL 197 1548 CG2 VAL 197 1549 C VAL 197 1550 O VAL 197 1551 N LEU 198 1552 CA LEU 198 1553 CB LEU 198	1514 NE1 TRP 192 17.691 1515 CZ2 TRP 192 18.525 1516 CZ3 TRP 192 17.537 1517 CH2 TRP 192 18.396 1518 C TRP 192 16.713 1519 O TRP 192 17.947 1520 N SER 193 15.991 1521 CA SER 193 16.676 1522 CB SER 193 15.672 1523 OG SER 193 14.658 1524 C SER 193 17.523 1525 O SER 193 18.582 1526 N CYS 194 17.064 1527 CA CYS 194 17.064 1527 CA CYS 194 17.886 1528 CB CYS 194 17.136 1529 SG CYS 194 15.813 1530 C CYS 194 15.813 1530 C CYS 194 19.195 1531 O CYS 194 20.223 1532 N GLY 195 19.137 1533 CA GLY 195 20.324 1534 C GLY 195 20.324 1534 C GLY 195 21.311 1535 O GLY 195 22.529 1536 N ILE 196 20.792 1537 CA ILE 196 20.792 1538 CB ILE 196 20.896 1539 CG2 ILE 196 19.649 1540 CG1 ILE 196 22.970 1541 CD1 ILE 196 22.970 1542 C ILE 196 22.970 1543 O ILE 196 22.970 1544 N VAL 197 21.527 1545 CA VAL 197 22.054 1546 CB VAL 197 22.054 1549 C VAL 197 22.057 1549 C VAL 197 23.193 1550 O VAL 197 24.220 1551 N LEU 198 23.579	1514 NE1 TRP 192 17.691 -1.086 1515 CZ2 TRP 192 18.525 -0.952 1516 CZ3 TRP 192 17.537 0.894 1517 CH2 TRP 192 18.396 -0.221 1518 C TRP 192 16.713 2.226 1519 O TRP 192 17.947 2.240 1520 N SER 193 15.991 1.706 1521 CA SER 193 16.676 1.118 1522 CB SER 193 15.672 0.467 1523 OG SER 193 14.658 1.368 1524 C SER 193 17.523 2.175 1525 O SER 193 18.582 1.866 1526 N CYS 194 17.064 3.420 1527 CA CYS 194 17.086 4.463 1528 CB CYS 194 17.136 5.799 1529 SG CYS 194 17.136 5.799 1529 SG CYS 194 15.813 5.829 1530 C CYS 194 19.195 4.643 1531 O CYS 194 19.195 4.643 1531 O CYS 194 19.195 4.643 1533 CA GLY 195 19.137 4.424 1533 CA GLY 195 20.324 4.541 1534 C GLY 195 20.324 4.541 1535 O GLY 195 22.529 3.624 1536 N ILE 196 20.792 2.225 1537 CA ILE 196 20.792 2.225 1537 CA ILE 196 20.896 -0.240 1539 CG2 ILE 196 20.896 -0.240 1539 CG2 ILE 196 20.896 -0.240 1539 CG2 ILE 196 22.970 -1.620 1541 CD1 ILE 196 22.970 -1.620 1542 C ILE 196 22.970 -1.620 1544 N VAL 197 22.054 2.257 1546 CB VAL 197 22.054 2.257 1546 CB VAL 197 22.054 2.257 1547 CG1 VAL 197 22.054 2.257 1548 CG2 VAL 197 19.986 1.740 1549 C VAL 197 23.193 3.270 1550 O VAL 197 24.220 3.168 1551 N LEU 198 23.026 4.231 1552 CA LEU 198 23.026 4.231	1514 NE1 TRP 192 17.691 -1.086 34.846 1515 CZ2 TRP 192 18.525 -0.952 37.215 1516 CZ3 TRP 192 17.537 0.894 38.439 1517 CH2 TRP 192 18.396 -0.221 38.368 1518 C TRP 192 16.713 2.226 32.364 1519 O TRP 192 17.947 2.240 32.345 1520 N SER 193 15.991 1.706 31.373 1521 CA SER 193 16.676 1.118 30.221 1522 CB SER 193 15.672 0.467 29.263 1523 OG SER 193 14.658 1.368 28.864 1524 C SER 193 17.523 2.175 29.506 1525 O SER 193 17.523 2.175 29.506 1526 N CYS 194 17.064 3.420 29.471 1527 CA CYS 194 17.064 3.420 29.471 1527 CA CYS 194 17.886 4.463 28.840 1528 CB CYS 194 17.136 5.799 28.793 1529 SG CYS 194 15.813 5.829 27.558 1530 C CYS 194 19.195 4.643 29.624 1531 O CYS 194 20.223 4.970 29.050 1532 N GLY 195 20.324 4.541 31.776 1534 C GLY 195 20.324 4.541 31.7480 1538 CB ILE 196 20.792 2.225 31.223 1537 CA ILE 196 20.792 2.225 31.223 1539 CG2 ILE 196 19.649 -0.143 30.132 1540 CG1 ILE 196 21.673 1.100 30.899 1538 CB ILE 196 20.896 -0.240 30.942 1539 CG2 ILE 196 21.673 1.300 30.415 1541 CD1 ILE 196 22.970 1.620 31.237 1542 C ILE 196 22.970 1.620 31.237 1543 O ILE 196 22.997 1.345 29.516 1543 O ILE 196 22.997 1.345 29.516 1543 O ILE 196 22.997 1.620 31.237 1544 N VAL 197 21.527 1.941 28.603 1545 CA VAL 197 22.054 2.257 27.278 1546 CB VAL 197 22.054 2.257 27.278 1548 CG2 VAL 197 21.593 3.495 25.106 1548 CG2 VAL 197 21.593 3.495 25.106 1551 N LEU 198 23.026 4.231 28.344 1552 CA LEU 198 23.026 4.231 28.344 1552 CA LEU 198 23.026 4.231 28.344

FIG.11A-37

	MOTA	1555	CD1	LEU	198		23.945	8.469	30.718	1.00	15.92	
	MOTA	1556	CD2	LEU	198	٠.	25.243	8.000	28.625	1.00	14.52	
	ATOM	1557	C	LEU	198		25.313	4.560	29.110	1.00	14.88	
	MOTA	1558	0	LEU	198		26.436	4.864	28.702	1.00	14.46	
	MOTA	1559	N	THR	199		25.117	3.639	30.044	1.00	14.39	
	MOTA	1560	CA	THR	199		26.250	2.909	30.623	1.00	16.47	
	MOTA	1561	CB	THR	199		25.766	1.920	31.698	1.00	14.59	:
	MOTA	1562	0G1	THR	199	:	25.085	2.643	32.728	1.00	15.04	
	MOTA	1563	CG2	THR	199		26.947	1.174	32.321	1.00	13.58	
	MOTA	1564	C	THR	199		27.005	2.156	29.523	1.00	17.44	
	MOTA	1565	0	THR	199	1.	28.237	2.192	29.465	1.00	18.28	•
	MOTA	1566	N	ALA	200		26.261	1.486	28.646	1.00	15.89	
	ATOM .	1567	CA	ALA	200		26.866	0.736	27.546	1.00	16.21	•
	MOTA	1568	CB	ALA	200		25.777	0.003	26.749	1.00	14.29	i, i.e.
	MOTA	1569	C	ALA	200	1722	27.662	1.660	26.623	1.00	17.57	
	MOTA	1570	0	ALA	200	ž H	28.781	1.337	26.225	1.00	18.68	
	MOTA	1571	N	MET			27.090		26.271		16.39	14.13
	MOTA	1572	CA	MET	201	• .	27.792	3.742	25.389		14.62	
	MOTA	1573	CB	MET			26.904		25.025		11.19	
	ATOM-	1574	CG	MET.	201	est 5	25.656		24.221			
	ATOM	1575	SD	MET		• • • • • • • • • • • • • • • • • • • •		6.071			18.10	
	ATOM	1576	CE	MET					24.918		13.57	*-
	ATOM	1577		MET	201			4.275			15.76	
	MOTA	1578			201		30.055	· .	25.296		15.99	
	ATOM	1579		LEU			29.086		27.325		15.81	
	MOTA	1580		LEU	202		30.258	•	28.014	1.00	17.08	*****
	MOTA	•	CB	LEU	202		29.805		29.195			
	ATOM	1582	CG	LEU			29.018	~	28.828		15.95	
•	ATOM	1583		LEU		٠.	28.622		30.095		12.72	
	MOTA	1584			202		29.870		27.910		16.64	
	MOTA	1585		LEU	202		31.309		28.512		18.70	
	MOTA	1586		LEU	202		32.440		28.815			
	MOTA	1587	N	ALA	203	•	30.956		28.592		17.89	
	ATOM	1588	CA	ALA	203		31.906		29.088		19.09	
	MOTA	1589	CB	ALA	203		31.509		30.493		16.50	
	ATOM	1590	C	ALA	203	٠	32.064		28.191	1.00		
	MOTA	1591	0	ALA	203	•	32.957				19.86	
	MOTA	1592	N	GLY	204		31.197		27.195		17.84	
	ATOM	1593		GLY	204		31.279		26.283		19.26	
	MOTA	1594	C	GLY	204		30.967		26.920		21.43	
	ATOM	1595	0	GLY	204		31.435				23.21	
	MOTA	1596	N	GLU	205		30.199	-2.074	28.002	1.00	20.75	

FIG.11A-38

1597	CA G	LU 205		29.806	-3.302	28.688	1.00 20.39
1598	CB G	LU 205		30.935	-3.826	29.588	1.00 22.16
1599	CG G	LU 205		31.143	-3.074	30.887	1.00 27.49
1600	CD G	LU 205		32.247	-3.681	31.751	1.00 29.22
1601	OE1 G	LU 205		32.138	-4.860	32.146	1.00 35.45
1602	0E2 G	LU 205		33.225	-2.971	32.040	1.00 28.80
1603	C G	LU 205		28.563	-3.054	29.518	1.00 18.62
1604	0 0	LU 205		28.305	-1.932	29.958	1.00 19.35
1605	N L	EU 206		27.779			1.00 19.99
1606				26.562	-4.013		1.00 20.37
1607	CB L	EU 206	•	25.543	-5.044		1.00 18.33
1608	CG L	EU 206		24.899	-4.783	28.643	1.00 20.09
1609	CD1 L	EU 206		25.952	-4.511		1.00 30.30 -
1610	CD2 L	_EU 206		24.075	-5.987		1.00 18.48
1611	C	EU 206		26.976	-4.290	31.944	1.00 21.02
1612	0. 1	LEU 206		27.769	-5.195	32.205	1.00 21.65
1613	N F	PRO 207		26.449	-3.510	32.898	1.00 21.06
1614	CD I	PRO 207	.	25.507	-2.400	32.678	1.00 18.20
1615	CA I	PRO 207	1	26.760	-3.646	34.323	1.00 21.75
1616	CB I	PRO 207	•	26.118	-2.405	34.932	1.00 19.82
1617	CG I	PRO 207	,	24.920	-2.200	34.055	1.00 17.27
1618	C	PRO 207	•	26.330	-4.929	35.027	1.00 23.19
1619	0 1	PRO 207	•	27.002	-5.363	35.958	1.00 25.40
1620	N	TRP 208		25.222	-5.533	34.600	1.00 20.85
1621	CA	TRP 208	}	24.759	-6.768	35.227	1.00 19.87
1622	CB	TRP 208	}	24.037	-6.449	36.542	1.00 17.82
1623	CG '	TRP 208	}	23.079	-5.294	36.431	1.00 16.93
1624	CD2	TRP 208	}	23.259	-3.986	36.978	1.00 15.33
1625	CE2	TRP 208	}	22.156	-3.203	36.564	1.00 19.36
1626	CE3	TRP 208	}	24.245	-3.394	37.777	1.00 16.72
1627	CD1	TRP 208	} %	21.906	-5.261	35.730	1.00 19.54
1628	NE1	TRP 208	3	21.344	-4.005	35.805	1.00 19.05
1629	CZ2	TRP 208	} · ·	22.017	-1.861	36.923	1.00 17.14
1630	CZ3	TRP 208	3	24.102	-2.057	38.137	1.00 17.80
1631	CH2	TRP 208	3	22.994	-1.306	37.708	1.00 17.63
1632	C.	TRP 208	3	23.847	·7.604	34.334	1.00 21.19
1633	0	TRP 208	3	23.243	-7.094	33.389	1.00 21.45
1634	N .	ASP 209)	23.758	-8.896	34.635	1.00 22.55
1635	CA ·	ASP 209	•			33.865	1.00 23.07
1636	CB ·	ASP 209)			34.317	1.00 24.77
1637	CG	ASP 209	•	24.456	-11.812		1.00 29.47
1638	OD1	ASP 209	•	24.996	-11.464	32.901	1.00 32.08
	1599 1600 1601 1602 1603 1604 1605 1606 1607 1608 1609 1610 1611 1612 1613 1614 1615 1616 1617 1618 1620 1621 1622 1623 1624 1625 1626 1627 1628 1629 1630 1631 1632 1633 1634 1635 1636 1637	1598 CB G 1599 CG G 1600 CD G 1601 OE1 G 1602 OE2 G 1603 C G 1604 O G 1605 N G 1606 CA G 1607 CB G 1609 CD1 G 1610 CD2 G 1611 C G 1612 O G 1613 N G 1614 CD G 1615 CA G 1616 CB G 1617 CG G 1618 C G 1619 O G 1620 N G 1621 CA G 1622 CB G 1624 CD2 1625 CE2 1626 CE3 1627 CD1 1628 NE1 1629 CZ2 1630 CZ3 1631 CH2 1632 C G 1634 N G 1635 CA G 1636 CB G 1637 CG	1598 CB GLU 205 1599 CG GLU 205 1600 CD GLU 205 1601 OE1 GLU 205 1602 OE2 GLU 205 1603 C GLU 205 1604 O GLU 205 1605 N LEU 206 1606 CA LEU 206 1607 CB LEU 206 1608 CG LEU 206 1609 CD1 LEU 206 1610 CD2 LEU 206 1611 C LEU 206 1613 N PRO 207 1614 CD PRO 207 1615 CA PRO 207 1616 CB PRO 207 1617 CG PRO 207 1618 C PRO 207 1619 O PRO 207 1619 CPRO 207 1620 N TRP 208 1621 CA TRP 208 1622 CB TRP 208 1623 CG TRP 208 1624 CD2 TRP 208 1625 CE2 TRP 208 1626 CE3 TRP 208 1627 CD1 TRP 208 1628 NE1 TRP 208 1629 CZ2 TRP 208 1631 CH2 TRP 208 1631 CH2 TRP 208 1632 C TRP 208 1633 O TRP 208 1634 N ASP 208 1635 CA ASP 208 1636 CB ASP 208 1637 CG ASP 208	1598 CB GLU 205 1599 CG GLU 205 1600 CD GLU 205 1601 OE1 GLU 205 1602 OE2 GLU 205 1603 C GLU 205 1604 O GLU 205 1605 N LEU 206 1606 CA LEU 206 1607 CB LEU 206 1608 CG LEU 206 1609 CD1 LEU 206 1610 CD2 LEU 206 1611 C LEU 206 1612 O LEU 206 1613 N PRO 207 1614 CD PRO 207 1615 CA PRO 207 1616 CB PRO 207 1616 CB PRO 207 1617 CG PRO 207 1618 C PRO 207 1619 O PRO 207 1619 O PRO 207 1620 N TRP 208 1621 CA TRP 208 1622 CB TRP 208 1624 CD2 TRP 208 1625 CE2 TRP 208 1626 CE3 TRP 208 1627 CD1 TRP 208 1628 NE1 TRP 208 1629 CZ2 TRP 208 1631 CH2 TRP 208 1632 C TRP 208 1633 O TRP 208 1631 CH2 TRP 208 1633 O TRP 208 1634 N ASP 209 1636 CB ASP 209	1598 CB GLU 205 30.935 1599 CG GLU 205 31.143 1600 CD GLU 205 32.247 1601 DE1 GLU 205 32.138 1602 DE2 GLU 205 33.225 1603 C GLU 205 28.563 1604 O GLU 205 28.305 1605 N LEU 206 27.779 1606 CA LEU 206 26.562 1607 CB LEU 206 25.543 1608 CG LEU 206 25.952 1610 CD2 LEU 206 24.899 1609 CD1 LEU 206 25.952 1610 CD2 LEU 206 24.075 1611 C LEU 206 26.976 1612 O LEU 206 26.7769 1613 N PRO 207 26.449 1614 CD PRO 207 26.760 1616 CB PRO 207 26.760 1616 CB PRO 207 26.760 1617 CG PRO 207 26.330 1619 O PRO 207 26.330 1619 O PRO 207 27.002 1620 N TRP 208 25.222 1621 CA TRP 208 24.037 1622 CB TRP 208 24.037 1623 CG TRP 208 23.079 1624 CD2 TRP 208 23.079 1625 CE2 TRP 208 23.259 1626 CE3 TRP 208 24.245 1627 CD1 TRP 208 24.245 1628 NE1 TRP 208 24.245 1629 CZ2 TRP 208 22.017 1630 CZ3 TRP 208 24.102 1631 CH2 TRP 208 23.259 1632 C TRP 208 24.945 1633 O TRP 208 23.259 1633 C TRP 208 24.945 1634 N ASP 209 23.758 1635 CA ASP 209 23.758 1636 CB ASP 209 23.087 1637 CG ASP 209 23.087	1598 CB GLU 205 30.935 -3.826 1599 CG GLU 205 31.143 -3.074 1600 CD GLU 205 32.247 -3.681 1601 0E1 GLU 205 32.138 -4.860 1602 0E2 GLU 205 33.225 -2.971 1603 C GLU 205 28.563 -3.054 1604 0 GLU 205 28.305 -1.932 1605 N LEU 206 26.562 -4.013 1607 CB LEU 206 26.562 -4.013 1607 CB LEU 206 25.543 -5.044 1608 CG LEU 206 25.952 -4.511 1610 CD2 LEU 206 26.952 -4.511 1610 CD2 LEU 206 26.976 -4.290 1611 C LEU 206 26.7769 -5.195 1613 N PRO 207 26.449 -3.510 1614 CD PRO 207 26.449 -3.510 1615 CA PRO 207 26.760 -3.646 1616 CB PRO 207 26.330 -4.929 1617 CG PRO 207 26.330 -4.929 1618 C PRO 207 26.330 -4.929 1619 O PRO 207 26.330 -4.929 1619 C PRO 207 27.002 -5.363 1620 N TRP 208 25.222 -5.533 1621 CA TRP 208 24.037 -6.449 1623 CG TRP 208 23.259 -3.986 1625 CE2 TRP 208 24.245 -3.394 1627 CD1 TRP 208 21.906 -5.261 1628 NE1 TRP 208 21.906 -5.261 1630 CZ3 TRP 208 22.156 -3.203 1626 CE3 TRP 208 24.245 -3.394 1627 CD1 TRP 208 22.156 -3.203 1628 NE1 TRP 208 22.156 -3.203 1629 CZZ TRP 208 22.156 -3.203 1620 CZ3 TRP 208 22.156 -3.203 1621 CA TRP 208 22.156 -3.203 1622 CB TRP 208 22.156 -3.203 1623 CG TRP 208 22.994 -1.306 1633 O TRP 208 23.243 -7.094 1634 N ASP 209 23.758 -8.896 1635 CA ASP 209 23.087 -11.256 1637 CG ASP 209 23.087 -11.256	1598 CB GLU 205 30.935 -3.826 29.588 1599 CG GLU 205 31.143 -3.074 30.887 1600 CD GLU 205 32.247 -3.681 31.751 1601 DE1 GLU 205 32.138 -4.860 32.146 1602 DE2 GLU 205 33.225 -2.971 32.040 1603 C GLU 205 28.563 -3.054 29.518 1604 O GLU 205 28.563 -3.054 29.518 1605 N LEU 206 27.779 -4.105 29.714 1606 CA LEU 206 26.562 -4.013 30.505 1607 CB LEU 206 26.562 -4.013 30.505 1607 CB LEU 206 25.543 -5.044 30.013 1608 CG LEU 206 25.952 -4.511 27.586 1610 CD2 LEU 206 24.075 -5.987 28.246 1611 C LEU 206 26.976 -4.290 31.944 1612 O LEU 206 27.769 -5.195 32.205 1613 N PRO 207 26.449 -3.510 32.898 1614 CD PRO 207 25.507 -2.400 32.678 1615 CA PRO 207 25.507 -2.400 32.678 1616 CB PRO 207 26.330 -4.929 35.027 1619 O PRO 207 26.330 -4.929 35.027 1619 O PRO 207 27.002 -5.363 35.958 1620 N TRP 208 25.222 -5.533 34.600 1621 CA TRP 208 24.037 -6.449 36.542 1623 CG TRP 208 24.037 -6.449 36.542 1624 CD2 TRP 208 24.037 -6.449 36.542 1625 CE2 TRP 208 23.079 -5.294 36.431 1626 CE3 TRP 208 24.245 -3.394 37.777 1627 CD1 TRP 208 21.344 -4.005 35.805 1630 CZ3 TRP 208 22.156 -3.203 36.564 1631 CH2 TRP 208 22.994 -1.306 37.708 1632 C TRP 208 22.994 -1.306 37.708 1633 O TRP 208 23.243 -7.094 33.389 1634 N ASP 209 22.901 -9.800 33.865 1636 CB ASP 209 22.901 -9.800 33.865 1636 CB ASP 209 22.901 -9.800 33.865

FIG.11A-39

MOTA	1639	OD2	ASP	209	24.981	-12.619	34.770	1.00 37.34
MOTA	1640	·C	ASP	209	21.439	-9.398	34.063	1.00 22.59
MOTA	1641	0	ASP	209	20.623	-9.498	33.143	1.00 21.49
MOTA	1642	N	GLN	210	21.123	-8.953	35.276	1.00 21.47
MOTA	1643	CA	GLN	210	19.775	-8.522	35.635	1.00 21.65
MOTA	1644	CB :	GLN	210	18.832	-9.729	35.725	1.00 21.36
MOTA	1645	CG	GLN	210	19.346	-10.845	36.622	1.00 22.91
MOTA	1646	CD	GLN	210	18.428	-12.051	36.635	1.00 24.69
MOTA	1647	0E1	GLN	210	18.188	-12.676	35.600	1.00 28.77
MOTA	1648	NE2	GLN	210	17.905	-12.383	37.810	1.00 29.76
MOTA	1649	:C	GLN	210			36.972	1.00 24.10
MOTA	1650	0	GLN	210	20.731		37.789	1.00 24.28
MOTA	1651	N	PRO .	211	18.874		37.214	
MOTA	1652	CD	PR0	211		-6.387		
ATOM	1653	CA	PR0	211			38.451	
MOTA	1654	CB	PR0	211				1.00 23.48
ATOM	1655	CG	PR0	211			37.072	
ATOM	1656	C	PR0	211		and the second second	39.634	3 .
ATOM	1657	0	PR0	211		The state of the s	40.127	
ATOM	1658	N	SER	212		-7.898	40.091	
ATOM	1659			212			41.214	
ATOM	1660	CB	SER	212			40.770	
ATOM	1661			212			39.621	1.00 37.91
ATOM	1662	C		212			42.349	
ATOM	1663	0		212	•		42.117	
ATOM	1664		ASP	213	18.834			1.00 35.30
ATOM		CA		213	-		44.741	
ATOM	1666			213			46.014	
ATOM	1667			213				1.00 52.63
MOTA	1668		ASP	213			47.204	
ATOM	1669			213				
ATOM	1670		ASP	213			44.632	
MOTA	1671		ASP	213		-10.070	45.180	1.00 41.19
MOTA	1672	N	SER	214			•	
ATOM	1673			214			43.721	•
MOTA	1674		SER	214			43.157	
MOTA	1675		SER	214			41.917	· · · · · · · · · · · · · · · · · · ·
MOTA	1676		SER	214		-12.063		
MOTA	1677		SER	214		-12.819		
MOTA	1678		CYS	215		-10.991		
ATOM	1679			215			41.045	
MOTA	1680	CB	CYS	215	22.557	-9.851	39.871	1.00 33.69

FIG.11A-40

1	MOTA	1681	SG	CYS	215	23.706	-9.553	38.523	1.00 33.80
,	ATOM	1682	С	CYS	215	24.223	-9.792	41.749	1.00 30.56
,	ATOM -	1683	0	CYS	215	23.976	-8.648	42.123	1.00 30.45
	MOTA	1684	N	GLN	216	25.410	-10.369	41.918	1.00 30.39
	MOTA	1685	CA	GLN	216	26.497	-9.687	42.602	1.00 27.76
	MOTA	1686	CB	GLN .	216	27.753	-10.569	42.621	1.00 28.61
	MOTA	1687	CG	GLN .	216	28.854	-10.012	43.510	1.00 32.96
	ATOM -	1688	CD	GLN	216	28.421	-9.895	44.963	1.00 42.47
	MOTA	1689	OE1	GLN:	216	28.866	-9.004	45.686	1.00 41.12
	MOTA	1690	NE2	GLN	216			45.398	1.00 48.90
	ATOM	1691	C	GLN	216	26.838	-8.319	42.014	1.00 26.11
	ATOM	1692	0	GLN	216	27.078	-7.375	42.759	1.00 23.93
	ATOM	1693	N	GLU	217	26.861	-8.212	40.688	1.00 25.76
	ATOM	1694	CA	GLU	217	27.176	-6.937	40.045	1.00 23.96
	MOTA	1695	CB	GLU	217	27.213	-7.092	38.521	1.00 24.39
	ATOM	1696	CĞ	GLU	217	28.404	-7.884	37.980	1.00 27.80
	MOTA	1697	CD	GLU	217	28.416	-9.327	38.453	1.00 30.02
	ATOM	1698	OE1	GLU	217	27.330	-9.944	38.514	1.00 26,98
	ATOM	1699	0E2	GLU	217	29.515	-9.845	38.754	1.00 34.80
	MOTA	1700	C	GLU	217	26.154	-5.868	40.432	1.00 22.37
	MOTA	1701	0	GLU	217	26.507	-4.701	40.629	1.00 20.68
	ATOM	1702	N	TYR	218	24.888	-6.261	40.547	1.00 22.07
	ATOM	1703	CA	TYR	218	23.858	-5.303	40.927	1.00 22.82
	ATOM	1704	CB	TYR	218	22.454	-5.858	40.664	1.00 24.84
	ATOM	1705	CG	TYR	218	21.371	-4.831	40.920	1.00 26.40
	ATOM	1706	CD1	TYR	218	21.373	-3.611	40.245	1.00 22.79
	ATOM	1707	CE1	TYR	218	20.402	-2.644	40.496	1.00 21.41
	ATOM	1708	CD2	TYR	218	20.363	-5.062	41.856	1.00 27.38
	ATOM	1709	CE2	TYR	218	19.385	-4.101	42.114	1.00 24.51
	ATOM	1710	CZ	TYR	218	19.413	-2.896	41.433	1.00 21.04
	ATOM -	1711	OH	TYR	218	18.469	-1.929	41.702	1.00 23.17
	ATOM	1712	C	TYR	218	23.991	-4.917	42.397	1.00 22.65
	ATOM	1713	0	TYR	218	23.811	-3.754	42.750	1.00 23.66
	ATOM	1714	N	SER	219	24.302	-5.888	43.256	1.00 25.18
	ATOM	1715	CA	SER	219	24.470	-5.600	44.681	1.00 24.31
	ATOM	1716	CB	SER	219	24.737	-6.889	45.471	1.00 26.30
•	ATOM	1717	OG	SER	219	23.648	-7.782	45.364	1.00 36.64
	ATOM	1718	C	SER	219	25.629	-4.628	44.888	1.00 22.69
	ATOM	1719	0	SER	219	25.527	-3.697	45.688	1.00 22.21
	ATOM	1720	N	ASP	220	26.725	-4.853	44.168	1.00 24.43
	MOTA	1721	CA	ASP	220	27.904	-3.992	44.257	1.00 24.43
	ATOM	1722	CB	ASP	220	28.990	-4.469	43.288	1.00 23.90

FIG.11A-41

ATOM	1723	CG ASP	220		29.662	-5.759	43.742	1.00 29.00
ATOM	1724	OD1 ASP	220		30.451	-6.320	42.954	1.00 35.56
MOTA	1725	OD2 ASP	220		29.406	-6.205	44.881	1.00 33.13
MOTA	1726	C ASP	220		27.532	-2.545	43.935	1.00 24.18
MOTA	1727	O ASP	220		28.007	-1.613	44.584	1.00 23.82
MOTA	1728	N TRP	221		26.679	-2.360	42.930	1.00 22.56
MOTA	1729	CA TRP	221		26.247	-1.016	42.545	1.00 20.51
MOTA	1730	CB TRP	221		25.414	-1.090	41.256	1.00 19.85
MOTA	1731	CG TRP	221		24.672		40.909	1.00 20.17
MOTA	1732	CD2 TRP	221		25.238	1.408	40.434	1.00 20.07
ATOM:	1733	CE2 TRP	221		24.163	2.309	40.226	1.00 17.90
MOTA	1734	CE3 TRP	221	17. 1	26.542	1.841	40.165	1.00 17.56
MOTA	1735	CD1 TRP	221		23.322	0.378	40.972	1.00 18.44
MOTA	1736	NE1 TRP	221	1.	23.008	1.653	40.563	1.00 18.49
MOTA	1737	CZ2 TRP	221	. **	24.360	3.614	39.758	1.00 15.58
MOTA	1738	CZ3 TRP	221		26.738	3.141	39.701	1.00 18.36
MOTA	1739	CH2 TRP	221		25.650	4.012	39.501	1.00 17.00
MOTA	1740	C TRP	221		25.446	-0.356	43.667	1.00 21.95
MOTA	1741	O TRP	221		25.662	0.810	43.995	1.00 21.87
ATOM	1742	N LYS	222		24.521	-1.099	44.262	1.00 24.52
MOTA	1743	CA LYS	222	mer i.	23.721	-0.543	45.343	1.00 26.38
MOTA	1744	CB LYS	222	171	22.596	-1.505	45.726	1.00 27.07
MOTA	1745	CG LYS	222	· · · · · · ·	21.565	-1.698	44.618	1.00 24.09
MOTA	1746	CD LYS	222	. A	20.299	-2.376	45.123	1.00 27.22
MOTA	1747	CE LYS	222		20.538	-3.831	45.493	1.00 25.58
MOTA	1748	NZ LYS	222		19.279	-4.473	45.958	1.00 28.43
MOTA	1749	C LYS	222		24.601	-0.233	46.553	1.00 28.55
MOTA	1750	0 LYS	222		24.251	0.601	47.385	1.00 29.05
MOTA	1751	N GLU	223		25.750	-0.898	46.635	1.00 29.11
MOTA	1752	CA GLU	223		26.691	-0.674	47.730	1.00 31.70
MOTA	1753	CB GLU	223		27.482	-1.950	48.026	1.00 35.51
MOTA	1754	CG GLU	223		26.650	-3.085	48.592	1.00 49.01
MOTA	1755	CD GLU	223		27.485	-4.311	48.900	1.00 57.47
MOTA	1756	OE1 GLU	223		28.415	-4.205	49.726	1.00 62.72
MOTA	1757	OE2 GLU	223		27.214	-5.381	48.313	1.00 63.79
ATOM	1758	C GLU	223		27.658	0.455	47.381	1.00 32.76
MOTA	1759	O GLU	223	•	28.578	0.756	48.144	1.00 33.37
ATOM	1760	N LYS	224		27.446	1.068	46.219	1.00 32.14
MOTA	1761	CA LYS	224		28.272	2.178	45.745	1.00 33.92
MOTA	1762	CB LYS	224		28.229	3.338	46.750	1.00 38.46
MOTA	1763	CG LYS	224		26.913	4.109	46.777	1.00 46.23
MOTA	1764	CD LYS	224		25.775	3.286	47.359	1.00 56.20

FIG.11A-42

ATOM	1765	CE	LYS	224		25.974	3.040	48.848	1.00 61.78
ATOM	1766	NZ	LYS	224		25.995	4.315	49.618	1.00 65.83
ATOM	1767	C	LYS	224		29.729	1.830	45.440	1.00 34.18
MOTA	1768	0	LYS	224		30.615	2.673	45.573	1.00 34.19
ATOM	1769	N	LYS	225		29.978	0.597	45.016	1.00 33.64
MOTA	1770	CA	LYS	225		31.336	0.172	44.688	1.00 35.23
MOTA	1771	CB	LYS	225		31.453	-1.347	44.837	1.00 36.69
ATOM	1772	CG	LYS	225		31.093	-1.853	46.230	1.00 40.35
MOTA	1773	CD	LYS	225		31.044	-3.377	46.290	1.00 46.38
MOTA	1774	CE.	LYS	225	٠.	32.383	-4.004	45.943	1.00 52.69
MOTA	1775	NZ	LYS	225		32.346	-5.490	46.067	1.00 60.52
ATOM	1776	C	LYS	225		31.670	0.588	43.254	1.00 36.02
MOTA	1777	. 0	LYS	225		31.918	-0.255	42.391.	1.00 34.27
MOTA	1778	N	THR	226	?	31.684	1.895	43.010	1.00 37.19
MOTA	1779	CA	THR	226		31.957	2.424	41.678	1.00 38.34
MOTA	1780	CB	THR	226	•	31.516	3.902	41.571	1.00 38.25
MOTA	1781	0G1	THR	226		32.145	4.670	42.602	1.00 38.16
MOTA	1782	CG2	THR	226		30.005	4.011	41.714	1.00 32.35
MOTA	1783	C	THR	226		33.409	2.303	41.227	1.00 39.24
MOTA	1784	0	THR	226		33.757	2.710	40.118	1.00 38.72
MOTA	1785	N	TYR	227		34.257	1.745	42.084	1.00 40.18
MOTA	1786	CA	TYR	227		35.658	1.560	41.733	1.00 39.40
MOTA	1787	CB	TYR	227		36.521	1.474	42.998	
MOTA	1788	CG	TYR	227	٠	36.050			
MOTA	1789	CD1		227		36.283	-0.916	43.797	1.00 41.13
MOTA	1790		. TYR	227	•	35.832	-1.867	44.709	1.00 37.13
MOTA	1791		2 TYR	227		35.353	0.831	45.143	1.00 38.67
MOTA	1792		2 TYR	227		34.897		46.060	1.00 39.51
MOTA	1793		TYR	227		35.140	-1.456	45.837	1.00 39.20
ATOM	1794	OH	TYR	227		34.680	-2.387	46.738	
ATOM	1795	C	TYR	227	<u>.</u>	35.776	0.280	40.914	1.00 39.74
MOTA	1796			227					1.00 40.52
MOTA	1797		LEU	228	•	34.643	•	•	1.00 39.05
MOTA	1798		LEU	228	-,	34.590	-1.634		
ATOM	1799		LEU	228		33.447	•	· · · · · · · · · · · · · · · · · · ·	1.00 40.20
ATOM	1800	CG	LEU	228		33.661			
ATOM	1801		L LEU	228		32.410		42.217	
MOTA	1802		2 LEU	228		34.859		41.740	
MOTA	1803		LEU	228		34.442	-1.390	•	
MOTA	1804		LEU	228		33.843		38.033	
ATOM	1805		ASN	229		34.987			
ATOM	1806	CA	ASN	229		35.041	-2.348	36.235	1.00 42.82

FIG.11A-43

ATOM	1807	СВ	ASN	229	34.836	-3.784	35.733	1.00 47.56
ATOM	1808	CG	ASN	229	35.542	-4.046	34.413	1.00 53.15
ATOM	1809	OD1		229	36.739	-3.789	34.276	1.00 50.92
ATOM	1810	ND2		229	34.806	-4.567	33.438	1.00 57.87
ATOM	1811	C	ASN	229	34.192	-1.399	35.389	1.00 41.75
ATOM	1812	Ō	ASN	229	34.726	-0.466	34.785	1.00 44.90
ATOM	1813	N	PRO	230	32.866	-1.608		1.00 37.73
ATOM	1814	CD	PRO	230		-2.470	36.114	1.00 33.11
ATOM	1815	CA	PRO	230	. 32.103		34.490	1.00 30.88
MOTA	1816	CB	PRO	230	30.680	-1.229	34.575	1.00 30.85
ATOM	1817	CG	PR0	230	30.624		35.958	1.00 30.64
ATON	1818	C	PR0	230	32.193	0.798	•	1.00 25.71
ATOM		0	PRO	230	32.654			1.00 24.73
ATOM	• •	N	TRP	231	31.782			1.00 23.16
ATON		CA	TRP	231	31.757		15	1.00 21.32
ATON	1822	•	TRP	231	31.099		38.020	
ATOM	1823	CG	TRP	231	29.965	/	38.087	
ATOM	1824		TRP		28.741		• •	
ATOM	1825		TRP	231	28.023		37.637	
ATOM		CE3		231	28.188		36.450	
ATOM	1827	2.3		231	the second secon	•	38.789	
MOTA	1828		TRP	231	28.773	•	38.523	
MOTA	1829	CZ2	TRP	231	26.774		37.071	1.00 19.20
•	1830		TRP	231		7.1	35.886	
ATOM	1831		TRP	231	26.255			
ATOM	1832	C	TRP	231	33.098	3.210	36.685	1.00 22.69
ATOM .	1833	0	TRP	231	33.138		36.503	1.00 20.98
MOTA	1834	N	LYS	232 ·	34.199	2.507	36.921	1.00 23.86
MOTA	1835	CA	LYS	232	35.487	3.199	36.992	1.00 25.79
MOTA	1836	CB	LYS	232	36.560	2.276	37.586	1.00 24.84
MOTA	1837	CG	LYS	232	36.812	0.989	36.824	1.00 33.96
MOTA	1838	CD	LYS	232	37.851	0.136	37.560	1.00 39.70
MOTA	1839	CE	LYS	232	38.112	-1.185	36.856	1.00 44.39
MOTA	1840	NZ	LYS	232	39.067	-2.042	37.620	1.00 48.06
ATOM	1841	C	LYS	232	35.962	3.760	35.649	1.00 25.77
MOTA	1842	0	LYS	232	36.920	4.530	35.596	1.00 26.85
MOTA	1843	N	LYS	· 233	35.277	3.393	34.570	1.00 24.28
MOTA	1844	CA	LYS	233	35.638	3.852	33.228	1.00 21.58
MOTA	1845	CB	LYS	233	35.460	2.714	32.220	1.00 21.63
ATOM	1846	CG	LYS	233	36.298			
MOTA	1847	CD	LYS	233	36.181			1.00 21.02
MOTA	1848	CE	LYS	233	34.839			

FIG.11A-44

MOTA	1849	NZ	LYS	233	34.817	-1.324	30.311	1.00 24.83
MOTA	1850	С	LYS	233	34.800	5.025	32.731	1.00 22.49
MOTA	1851	0	LYS	233	35.041	5.545	31.642	1.00 22.51
MOTA	1852	N	ILE .	234	33.848	5.471	33.533	1.00 23.05
MOTA	1853	CA	ILE	234	32.933	6.504	33.062	1.00 23.85
ATOM	1854	CB	ILE	234	31.526	6.124	33.584	1.00 18.85
ATOM	1855	CG2	ILE	234	30.523	7.242	33.345	1.00 16.49
MOTA	1856	CG1	ILE	234	31.128	4.813	32.893	1.00 16.83
MOTA	1857	CD1	ILE -	234	29.773	4.256	33.265	1.00 15.87
MOTA	1858	C	ILE	234	33.206	8.015	33.175	1.00 26.97
ATOM	1859	0	ILE	234	33.655	8.629	32.202	1.00 34.42
ATOM	1860	N.	ASP	235	32.953	8.592	34.339	1.00 25.59
ATOM	1861	CA	ASP	235	33.136	10.025	34.646	.1.00 24.80
MOTA	1862	CB	ASP	235	32.528	10.995	33.623	1.00 22.82
ATOM	1863	CG	ASP	235	33.342	12.289	33.509	1.00 29.58
ATOM	1864	QD1	ASP	235	33.015	13.294	34.187	1.00 28.74
MOTA	1865	OD2	ASP	235	34.341	12.292	32.758	1.00 24.88
ATOM	1866	C	ASP:	235	32.512	10.282	35.990	1.00 22.84
MOTA	1867	0	ASP	235	31.766	9.448	36.503	1.00 19.57
ATOM	1868	N	SER	236	32.824	11.450	36.540	1.00 22.97
MOTA	1869	CA	SER	236	32.144	11.667	37.793	1.00 23.58
MOTA	1870	CB	SER	236	32.929	12.818	38.441	1.00 22.04
ATOM	1871	OG	SER	236	32.992	13.941	37.583	1.00 27.67
MOTA	1872	C	SER	236	30.991	12.390	- 37.096	1.00 21.73
MOTA	1873	0	SER	236	29.952	12.112	37.692	1.00 19.53
MOTA	1874	N	ALA	237	31.031	13.258	36.083	1.00 20.61
MOTA	1875	CA	ALA	237	29.816	13.944	35.639	1.00 19.05
MOTA	1876	CB	ALA	237	30.137	14.988	34.566	1.00 14.22
MOTA	1877	C	ALA	237	28.746	12.973	35.134	1.00 17.61
MOTA	1878	0	ALA	237	27.639	12.950	35.664	1.00 17.49
MOTA	1879	N	PRO	238	29.049	12.175	34.097	1.00 15.48
MOTA	1880	CD	PRO	238	30.217	12.121	33.199	1.00 15.00
MOTA	1881	CA.	PRO	238	27.999	11.252	33.646	1.00 16.82
MOTA	1882	CB	PRO	238	28.572	10.670		1.00 13.41
MOTA	1883	CG	PRO	238	30.067	10.766	32.552	1.00 10.20
MOTA	1884	C	PRO	238	27.670	10.183	34.694	1.00 14.91
MOTA	1885	0	PRO	238	26.539	9.701	34.770	1.00 14.08
MOTA	1886	N -	LEU	239	28.657	9.815	35.508	1.00 16.50
MOTA	1887	CA	LEU	239	28.434	8.819	36.554	1.00 17.72
MOTA	1888	CB	LEU	239	29.744	8.522	37.296	1.00 18.57
MOTA	1889	CG	LEU	239	30.096	7.069	37.643	1.00 22.40
MOTA	1890	CD1	LEU	239	31.090	7.086	38.795	1.00 23.81

FIG.11A-45

				· · · · · · · · · · · · · · · · · · ·						
MOTA	1891	CD2	LEU	239		28.873	6.257	38.017	1.00	22.04
MOTA	1892	С	LEU	239		27.394	9.351	37.543	1.00	17.57
MOTA	1893	0 "	LEU	239		26.543	8.605	38.026	1.00	16.91
MOTA	1894	N	ALA	240		27.464	10.645	37.846	1.00	16.87
MOTA	1895	CA	ALA	240	•	26.508	11.254	38.766	1.00	17.94
MOTA	1896	·CB	ALA	240		26.867	12.725	39.024	1.00	15.97
MOTA	1897	.C	ALA	240		25.091	11.143	38.198	1.00	16.55
MOTA	1898	0	ALA	240		24.136	10.974	38.950	1.00	16.22
ATOM	1899	N	LEU	241		24.954	11.241	36.878	1.00	15.65
MOTA	1900	CA	LEU	241	·.	23.627	11.111	36.264	1.00	15.31
MOTA	1901	CB	LEU	241		23.652	11.540	34.785	1.00	12.35
MOTA	1902	CG	LEU	241	'	22.354	11.270	33.991	1.00	13.16
ATOM	1903	CD1	LEU	241	<i>.</i> / .	21.170	11.991	34,606	1.00	14.35
MOTA	1904	CD2	LEU	241		22.535	11.720	32.557	1.00	12.21
MOTA	1905	C	LEU	241	14.5°	23.175	9.655	36.384		16.01
ATOM	1906	0	LEU	241		22.025	9.377	36.739	1.00	15.78
MOTA	1907	N	LEU.	242		24.076	·	36.095		15.89
MOTA	1908	CA	LEU	242		23.734		36.194		16.35
MOTA	1909	CB	LEU	242	· · · · · · · · · · · · · · · · · · ·	24.942	6.430	35.808		17.35
ATOM	1910	CG	LEU	242	84.1 L	25.054		34.500		22.28
ATOM			LEU	242				33.505		16.11
MOTA	1912	CD2	LEU	242				33.896		5
MOTA	1913	C	LEU	242				37.624		16.88
ATOM	1914	0	LEU	242	11 11 11	22.418		37.834		15.72
MOTA	1915	. N. A.				23.883		38.609		18.68
ATOM				243		23.507		40.004		17.89
MOTA			HIS	243	." -			40.958		17.93
MOTA		·	HIS		 	25.587		41.458		27.59
MOTA			HIS	243		25.622		42.232		27.35
MOTA			l HIS	243		26.900		41.176		30.65
MOTA				243		27.695		41.755		31.26
MOTA			2 HIS	243		26.944		42.402	•	27.77
MOTA			HIS	243		22.069		40.265		17.20
MOTA			HIS	243		21.425		41.189		17.65
MOTA			LYS			21.577		39.460		17.08
MOTA				244		20.212		39.617		17.67
ATOM			LYS			20.137	10.751			15.78
MOTA			LYS	_		20.904	11.670	40.163		19.56
MOTA			LYS			20.750	13.143	39.815		:19.62
ATOM			LYS			21.549	13.543			15.60
ATOM			LYS			21.582	15.043			18.38
ATOM	1932	С	LYS	244		19.213	8.447	38.805	1.00	16.94

FIG.11A-46

								·····
ATOM	1933	0	LYS	244	18.044	8.339	39.170	1.00 16.00
MOTA	1934	N	ILE	245	19.681	7.849	37.713	1.00 15.92
MOTA	1935	CA	ILE	245	18.812	7.023	36.871	1.00 14.11
MOTA	1936	CB	ILE	245	19.338	6.955	35.404	1.00 12.65
MOTA	1937	CG2	ILE	245	18.465	5.982	34.573	1.00 14.27
MOTA	1938	CG1	ILE	245	19.307	8.352	34.776	1.00 11.69
ATOM	1939	CD1	ILE	245	19.923	8.441	33.379	1.00 17.04
MOTA	1940	C	ILE	245	18.684	5.586	37.387	1.00 13.68
MOTA	1941	0 .	ILE	245	17.583	5.041	37.481	1.00 15.46
ATOM	1942	N	LEU	246	19.809	4.963	37.724	1.00 14.19
MOTA	1943	CA	LEU	246	19.765	3.574	38.173	1.00 15.34
MOTA	1944	CB	LEU	246	21.062	2.865	37.776	1.00 13.75
MOTA	1945	CG	LEU	246	21.346	2.984	-t36_278	1.00 10.86
ATOM	1946	CD1	LEU	246	22.703	2.364	35.923	1.00 12.91
ATOM	1947	CD2		246	20.211	2.303	35.512	1.00 14.82
MOTA	1948	C.	LEU	246	19.477	3.429	39.663	1.00 18.00
ATOM	1949	0	LEU	246	20.229	2.803	40.422	1.00 18.52
MOTA	1950	N	VAL	247	18.357	4.022	40.057	1.00 17.55
ATOM	1951	CA	VAL	247	17.881	4.000	41.433	1.00 16.51
MOTA	1952	CB	VAL	247	17.268	5.356	41.795	1.00 15.11
MOTA	1953		VAL	247	16.553	5.278	43.136	1.00 19.34
MOTA	1954		VAL	247	18.380	6.408	41.842	1.00 16.66
MOTA	1955	C	VAL	247	16.834	2.899	41.513	1.00 18.51
ATOM	1956	.0	VAL	247	15.903		40.709	1.00 18.14
ATOM		N	GLU	248	16.990	2.000	42.481	1.00 16.77
MOTA	1958	CA	GLU	248	16.078	0.864	42.613	1.00 17.92
ATOM	1959	CB	GLU	248	16.522	-0.034	43.767	1.00 19.95
ATOM	1960	CG	GLU	248	15.805	-1.376	43.799	1.00 28.58
ATOM	1961	CD	GLU	248	16.404	-2.315	44.822	1.00 43.56
ATOM	1962		GLU	248	16.396	-1.965	46.021	1.00 48.24
ATOM	1963		GLU	248	16.889	-3.396	44.425	1.00 46.86
ATOM	1964		GLU	248	14.605	1.224		1.00 18.01
MOTA	1965	0	GLU	248	13.741	0.633	42.131	1.00 17.60
ATOM	1966	N	ASN	249	14.317	2.185	43.652	1.00 17.72
MOTA	1967	CA	ASN	249	12.940	2.611	43.886	1.00 16.89
MOTA	1968	CB.	ASN	249	12.866	3.392	45.206	•
MOTA	1969	CG	ASN	249	11.480	3.970	45.480	1.00 21.33
ATOM	1970		ASN	249	10.562	3.832	44.676	
MOTA	1 971		ASN	249	11.331	4.624	46.631	1.00 17.12
ATOM	1972	C	ASN	249	12.480	3.483	42.716	1.00 15.11
MOTA	1973	0	ASN	249	12.954	4.607		1.00 16.33
MOTA	1974	N	PRO	250	11.550	2.978	41.880°	1.00 15.15

FIG.11A-47

MOTA	1975	CD	PR0	250	10.830	1.694	41.960	1.00 1	6.54	
MOŢA	1976	CA	PR0	250	11.080	3.774	40.737	1.00 1	5.50	•
ATOM	1977	CB	PRO	250	10.110	2.825	40.025	1.00 1	4.37	
ATOM	1978	CG	PRO	250	9.569	1.973	41.153	1.00	L3.99	
MOTA	1979	C	PRO	250	10.437	5.111	41.105	1.00 1	16.72	
ATOM	1980	0	PRO	250	10.409	6.039	40.298	1.00	L6.34	
ATOM	1981	a N	SER	251	9.910	5.211	42.321	1.00	17.93	
MOTA	1982	CA	SER	251	9.296	6.460	42.744	1.00	18.37	
MOTA	1983	CB	SER	251	8.391	6.212	43.954	1.00	18.23	
MOTA	1984	OG	SER	251	7.326	5.351	43.584	1.00 2	20.47	:
MOTA	1985	C	SER	251	10.347	7.524	43.060	1.00	18.31	
ATOM	1986	. 0	SER	251	10.075	8.720	42.944	1.00 2	21.19	
MOTA	1987	N ·	ALA	252	11.549	7.092	43.430	1.00	17.56	11.5
ATOM	1988	CA	ALA	252	12.638	8.020	43.749	1.00	16.14	$\mathcal{N}_{1}(M)$
MOTA	1989	CB	ALA	252	13.471	7.479	44.919	1.00	17.50	. #KT 1
MOTA	1990	C	ALA	252	13.545	8.257	42.544	1.00	16.13	talian en 1750 en 1860. Spear de la company
MOTA	1991	0	ALA	252	14.355	9.184	42.533		18.76	
MOTA	1992	N	ARG	253			41.530		16.12	
MOTA	Commercial	:		253			40.322		16.16	
MOTA	1994		ARG	253			39.382		15.22	7
ATOM	1995		ARG	253			38.149			
MOTA	1996		ARG	253			37.433		_	
ATOM	1997		ARG	253			38.391		15.13	
MOTA	1998		ARG	253	13.637		38.264		and the second second	
MOTA	1999			253	13.671		39.199		10.88	MADE.
MOTA	2000	NH2		253	12.849		37.203		13.34	
MOTA	2001			253	13.998		39.625	•	17.48	Sept 1
MOTA	2002		ARG	253	12.889		39.624	· · ·	16.77	N. 11.
MOTA	2003	•		254	15.054	•	39.033		15.92	
MOTA	2004		ILE	254	14.952		38.346		14.89	1,144,877
MOTA	2005		ILE	254	16.359				16.24	• .:
•	2006			-	16.867					
ATOM	2007			254	15.305		37.390		16.57	
ATOM	2008			254	17.679		37.079		15.04	•
MOTA	2009		ILE	254	13.981		37.164		16.39	
MOTA	2010	0	ILE	254					17.01	•
MOTA	2011		THR	255	13.242		36.908		17.02	
MOTA	2012	CA	THR	255	12.292	11.692	35.800			
ATOM	2013	CB	THR	255	11.037	12.517	36.147			
ATOM	2014		THR	255	11.433	13.837	36.542		19.30	•
MOTA	2015		THR	255	10.263					
MOTA	2015	C	THR	255	12.997	12.370	34.635	1.00	17.07	

FIG.11A-48

	ATOM	2017	0	THR	255	14.058	12.959	34.808	1.00 16.74
	ATOM	2018	N .	ILE	256	12.410	12.321	33.450	1.00 18.82
	MOTA	2019	CA	ILE	256	13.070	12.954	32.320	1.00 18.31
	MOTA	2020	CB	ILE	256	12.393	12.576	30.995	1.00 16.73
	MOTA	2021	CG2	ILE	256 .	13.076	13.305	29.844	1.00 16.91
	MOTA	2022	CG1	ILE	256	12.482	11.058	30.805	1.00 15.14
	MOTA	2023	CD1	ILE	256	11.814	10.538	29.555	1.00 17.21
	MOTA	2024	C	ILE	256	13.162	14.472	32.461	1.00 18.90
	MOTA	2025	0	ILE	256	14.182	15.062	32.112	1.00 19.82
	MOTA	2026	N	PRO	257	12.099	15.135	32.959	1.00 19.76
	MOTA	2027	CD	PRO	257	10.697	14.733	33.185	1.00 18.99
	MOTA	2028	CA	PRO	257	12.256	16.590	33.079	1.00 19.66
-	ATOM	2029	CB	PRO	257	10.948 -	17:.019	33.739	1.00 19.59
	ATOM	2030	CG	PRO	257	9.953	16.075	33.104	1.00 19.54
	MOTA	2031	C	PR0	257	13.494	16.949	33.911	1.00 19.08
	ATOM	2032	0	PRO	257	14.176	17.941	33.637	1.00 19.37
	ATOM	2033	N	ASP	258	13.794	16.133	34.917	1.00 19.06
	MOTA	2034	CA	ASP	258	14.958	16.373	35.760	1.00 18.81
	MOTA	2035	CB	ASP	258	14.735	15.728	37.128	1.00 18.17
	ATOM	2036	CG	ASP	258	13.772	16.542	37.978	1.00 23.21
	ATOM	2037	OD1	ASP	258	13.193	16.012	38.948	1.00 23.28
	MOTA	2038	0D2	ASP	258	13.611	17.738	37.652	1.00 23.96
	ATOM	2039	C	ASP	258	16.266	15.922	35.101	1.00 18.56
	ATOM	2040	0	ASP_	258	17.327	16.504	35.349	1.00 20.13
	ATOM	2041	N	ILE	259	16.197	14.906	34.246	1.00 18.38
	ATOM	2042	CA	ILE	259	17.392	14.471	33.531	1.00 19.58
	ATOM	2043	CB	ILE	259	17.114	13.239	32.618	1.00 16.93
	ATOM	2044	CG2	ILE	259	18.241	13.063	31.600	1.00 14.51
	ATOM	2045	CG1	ILE	259	16.994	11.966	33.464	1.00 16.72
	ATOM -	2046		ILE	259	16.489	10.748	32.677	1.00 11.48
	ATOM	2047	·.C	ILE	259	17.823	15.659	32.659	1.00 21.42
	MOTA	2048	0	ILE	259	19.005	15.958	32.543	1.00 20.64
	MOTA	2049	N	LYS	260	16.851	16.354	32.070	1.00 23.12
	MOTA	2050	CA	LYS	260	17.152	17.499	31.208	1.00 23.95
	MOTA	2051	CB	LYS	260	15.876	18.020	30.538	1.00 25.83
	MOTA	2052	CG	LYS	260	15.150	19.064	31.356	1.00 38.93
	ATOM	2053	CD	LYS	260	13.885	19.551	30.678	1.00 48.71
	ATOM	2054	CE	LYS	260	13.278	20.709	31.455	1:00 44.94
	MOTA	2055	NZ	LYS	260	14.210	21.872	31.510	1:00 42.57
	MOTA	2056	C	LYS	260	17.827	18.646	31.961	1.00 22.24
	MOTA	2057	0	LYS	260	18.369		31.340	1.00 22.70
	MOTA	2058	N	LYS	261	17.787			1.00 20.88

FIG.11A-49

								• "
ATOM	2059	CA	LYS	261	18.402	19.628	34.129	1.00 21.74
MOTA	2060	CB	LYS	261	17.474	19.984	35.298	1.00 24.31
MOTA	2061	CG	LYS	261	16.176	20.661	34.881	1.00 33.19
ATOM	2062	CD	LYS	261	15.245	20.857	36.071	1.00 47.75
MOTA	2063	CE	LYS	261	14.008	21.650	35.680	1.00 57.96
MOTA	2064	NZ	LYS	261	13.280	21.031	34.537	1.00 64.19
MOTA	2065	C	LYS	261	19.750	19.181	34.687	1.00 21.66
MOTA	2066	0	LYS	261	20.462	19.964	35.320	1.00 23.16
ATOM	2067	N	ASP.	262	20.105	17.926	34.442	1.00 19.55
MOTA	2068	·CA	ASP	· 262	21.352	17.371	34.950	1.00 19.62
ATOM	2069	CB	ASP	262	21.419	15.874	34.618	1.00 18.71
MOTA	2070	CG	ASP	262	22.781	15.266	34.903	1.00 15.55
MOTA	2071	OD1	ASP	262			33.955	
ATOM	2072	OD2					36.064	
MOTA	2073	C		262		18.102	34.437	
HOTA	2074			262			33.294	
	2075		ARG	263			35.290	1.00 18.70
MOTA	2076				24.825			1.00 18.98
MOTA	2077		ARG	263		18.937		1.00 20.52
	2078						35.820	1.00 28.61
MOTA		CD			27.963		37.068	1.00 38.40
MOTA	2080		ARG		28.937			
711011	2081		•		28.637		37.499	
MOTA	2082			263	27.375	•	37.665	1.00 58.45
MOTA				263	•		37.626	1.00 62.85
MOTA	2084		ARG				33.700	
MOTA			ARG		25.850		32.769	
MOTA			TRP	264			33.684	
MOTA			TRP	264	26.390		32.531	
MOTA	2088		TRP	264	26.684		32.788	
MOTA	2089			264	27.354		31.610	
ATOM -	•		TRP	264	26.733	13.407		1.00 16.57
MOTA	2091		TRP	264	27.715	13.090		1.00 14.53
MOTA	2092		TRP	264	25.437	12.878		1.00 15.64
MOTA	2093		TRP	264	28.652			1.00 12.93
MOTA	2094		TRP	264 264	28.875	13.722		1.00 13.60
MOTA	2095		TRP	264	27.446	12.269	28.570	1.00 14.44
MOTA	2096		TRP	264	25.168		29.391	1.00 13.46
MOTA	2097		TRP	264	26.166			
MOTA	2098		TRP	264 264	25.545			1.00 15.93
MOTA	2099		TRP		26.064			1.00 13.92
MOTA	2100	N	TYR	265	24.240	16.339	31.393	1:00 15.00

FIG.11A-50

MOTA	2101	CÂ	TYR	265	•	23.342	16.447	30.257	1.00 13.62
MOTA	2102	CB T	TYR	265		21.895	16.265	30.738	1.00 11.92
MOTA	2103	CG	TYR	265		20.888	16.112	29.629	1.00 15.19
MOTA	2104	CD1	TYR	265		20.259	17.220	29.060	1.00 16.33
MOTA	2105	CE1	TYR	265		19.317	17.062	28.039	1.00 16.52
MOTA	2106	CD2	TYR	265		20.555	14.843	29.150	1.00 13.45
MOTA	2107	CE2	TYR	265		19.628	14.676	28.148	1.00 13.56
MOTA	2108	CZ	TYR	265		19.010	15.781	27.594	1.00 16.67
MOTA	2109	OH	TYR	265		18.084	15.582	26.608	1.00 19.16
MOTA	2110	C .	TYR	265		23.508	17.798	29.551	1.00 14.32
MOTA	2111	0	TYR	265		23.459	17.882	28.322	1.00 15.01
MOTA	2112	N	ASN	266		23.751	18.847	30.335	1.00 15.47
MOTA	2113	CA	ASN	266		23.897	20.193	29.790	1.00 17.01
MOTA	2114	CB	ASN'	266		23.166	21.184	30.704	1.00 17.43
MOTA	2115	CG	ASN	266		21.661	21.021	30.636	1.00 19.60
MOTA	2116	OD1	ASN:	266	:	21.030	21.428	29.659	1.00 21.30
MOTA	2117	ND2		266		21.080	20.396	31.661	1.00 19.15
MOTA	2118	C	ASN	266		25.330	20.676	29.552	1.00 18.03
MOTA	2119	0	ASN	266		25.536	21.820	29.154	1.00 16.54
MOTA	2120	N	LYS	267		26.319	19.815	29.773	1.00 18.76
MOTA	2121	CA	LYS	267		27.716	20.221	29.574	1.00 18.99
MOTA	2122	CB	LYS	267		28.666	19.244	30.273	1.00 24.39
MOTA	2123	CG	LYS	267		28.804	19.442	31.769	1.00 35.68
MOTA		CD	LYS	267			18.424	32.368	1.00 48.27
MOTA	2125		LYS	267		31.138	18.481	31.702	1.00 51.14
MOTA	2126		LYS	267		31.800	19.802	31.888	1.00 56.24
MOTA	2127	C	LYS	267	•	28.123	20.307	28.110	1.00 19.25
ATOM	2128	Q	LYS	267		27.919	19.365	27.350	1.00 18.64
MOTA	2129	N	PRO	268		28.708	21.444	27.694	1.00 21.76
ATOM	2130	CD	PRO	268		28.826	22.742	28.378	1.00 22.13
MOTA	2131		PRO	268		29.119	21.547	26.289	1.00 23.47
MOTA	2132	CB	PRO	268		29.556			1.00 22.99
MOTA	2133		PRO	268		28.746			1.00 24.42
ATOM	2134	C	PRO	268	₹.		•	26.084	•
ATOM	2135	0	PRO	268		31.280			1.00 22.64
MOTA	2136	N	LEU	269		30.132		•	1.00 21.85
MOTA	2137	CA	LEU	269		31.155	18.667		1.00 23.57
MOTA	2138	CB	LEU	269		30.751			1.00 23.55
ATOM	2139	CG	LEU	269	•	30.576		26.982	1.00 22.37
MOTA	2140		LEU	269	•	29.980	15.818		1.00 23.62
MOTA	2141		LEU	269	•	31.920	17.367		1.00 21.81
MOTA	2142	С	LEU	269		31.442	18.424	23.394	1.00 26.56

FIG.11A-51

ATOM	2143	0	LEU	269	32.592	18.228	23.012	1.00 26.46
MOTA	2144	N	LYS	270	30.400	18.421	22.571	1.00 28.53
ATOM	2145	CA	LYS	270	30.595	18.128	21.158	1.00 31.73
MOTA	2146	CB	LYS	270	29.790	16.881	20.777	1.00 33.10
MOTA	2147	CG	LYS	270	30.179	16.292	19.431	1.00 37.08
MOTA	2148	CD	LYS	270	29.461	14.981	19.167	1.00 35.35
MOTA	2149	CE	LYS	270	29.881	14.383	17.833	1.00 33.40
MOTA	2150	NZ	LYS	270	29.137	13.123	17.546	1.00 40.91
MOTA	2151	C ·	LYS	270	30.290	19.241	20.171	1.00 34.07
MOTA	2152	0	LYS	270	29.304	19.968	20.301	1.00 34.39
MOTA	2153	N	LYS	271	31.162	19.358	19.177	1.00 36.37
MOTA	2154	CA	LYS	271	31.018	20.349	18.124	1.00 39.68
MOTA	2155	CB	LYS	271	32.345	20.528	17.381	1.00 40.41
MOTA	2156	CG	LYS	271	33.534	20.899	18.259	1.00 37.20
MOTA	2157	CD	LYS					1.00 32.85
MOTA	2158	CE		271				
MOTA	2159	NZ						1.00 21.18
MOTA	2160	C	LYS		29.975			
MOTA	2161	0						1.00 39.59
MOTA	2162	N		-				1.00 43.87
MOTA	2163					•	15.377	the second of th
						the second second second		1.00 49.32
MOTA	2165	0	GLY	272	30.317			1.00 49.55
	2166				28.306			1.00 50.80
MOTA	2167			273	28.850		12.240	
MOTA	2168				27.749		11.234	
ATOM	2169	•			29.998		11.547	·
ATOM	2170	0	-	273	30.024		11.501	
MOTA	2171			274	30.945		11.012	
, ti Oii	2172				32.101	· ·		1.00 57.69
MOTA			ALA	274	33.043	17.591	9.883	1.00 55.97
MOTA	2174			274	31.681			1.00 58.75
ATOM	2175		ALA	274	31.092	19.018		•
ATOM	2176		ALA	275	31.991	20.833		1.00 60.19
MOTA	2177			275	31.653			. 1.00 61.34
ATOM	2178			275	32.417		•	1.00 63.01
MOTA	2179	C.	ALA	275	30.155			1.00 62.70
MOTA	2180	0	ALA	275	29.687	21.161		1.00 64.42
MOTA	2181	N	ALA	276	29.406			1.00 63.05
MOTA	2182	CA		276	27.959			
MOTA	2183			276 °	27.300			1.00 65.25
MOTA	2184	C	ALA	276	27.409	23.722	9.302	1.00 66.43

FIG.11A-52

ATOM	2185 (CT1 ALA	276		26.726	24.582	8.707	1.00 66.01
MOTA	2186	OT ALA	276		27.665	23.761	10.524	1.00 72.06
MOTA	2187	OH2 WAT	500		7.288	0.582	30.446	1.00 12.93
MOTA	2188	OH2 WAT	501		7.551	-2.385	30.926	1.00 14.51
MOTA	2189	OH2 WAT	502		15.648	-3.549	26.581	1.00 12.66
MOTA	2190	OH2 WAT	503		22.995	-4.531	32.505	1.00 14.00
MOTA	2191	OH2 WAT	504		12.370	-2.139	29.668	1.00 12.75
ATOM	2192	OH2 WAT	505	•	8.243	1.795	37.412	1.00 13.95
MOTA	2193	OH2 WAT	506		12.211	-1.687	42.460	1.00 18.17
MOTA	2194	OH2 WAT	507		12.547	0.038	27.856	1.00 14.35
MOTA	2195	OH2 WAT	508		9.787	10.899	33.147	1.00 15.08
MOTA	2196	OH2 WAT	510		11.744	7.842	36.365	1.00 15.19
MOTA	2197	OH2 WAT	511		9.925	-3.492	29.777	1.00 15.10
MOTA	2198	OH2 WAT	512		9.590	8.537	34.696	1.00 17.43
MOTA	2199	OH2 WAT	513	:	2.021	3.295	33.836	1.00 15.34
MOTA	2200	OH2 WAT	514		6.563	13.229	27.860	1.00 18.19
MOTA	2201	OH2 WAT	515		10.555	8.269	38.785	1.00 18.00
MOTA	2202	OH2 WAT	516		10.674	15.405	22.497	1.00 19.56
MOTA	2203	OH2 WAT	517		25.750	15.101	36.287	1.00 17.00
MOTA	2204	OH2 WAT	518		4.386	6.182	34.218	1.00 15.43
MOTA	2205	OH2 WAT	519		13.712	-1.171	31.851	1.00 19.69
MOTA	2206	OH2 WAT	520		27.652	18.967	23.808	1.00 20.13
MOTA	2207	OH2 WAT	521		14.113	-4,152	28.944	1.00 16.61
MOTA	2208	OH2 WAT	522		8.101	9.135	38.813	
ATOM	2209	OH2 WAT	523		6.549	1.866	39.438	1.00 17.99
ATOM	2210	OH2 WAT	524		8.387	10.486	30.847	1.00 15.91
ATOM	2211	OH2 WAT	525		12.082	9.839	11.918	1.00 19.48
MOTA	2212	OH2 WAT	526		18.804	-3.707	34.246	1.00 13.10
ATOM	2213	OH2 WAT	527		13.250	13.468	39.304	1.00 19.10
MOTA	2214	OH2 WAT			7.275		36.188	1.00 19.69
ATOM	2215	OH2 WAT	529		5.361	7.284	36.859	1.00 17.02
MOTA		OH2 WAT	530		8.547		29.494	
MOTA		OH2 WAT	531		33.657			1.00 19.62
ATOM	2218	OH2 WAT	532		23.095	17.810	38.035	1.00 20.16
ATOM	2219		533		7.044	4.516		
ATOM	2220	OH2 WAT	534	•	8.572		21.497	
ATOM	2221	OH2 WAT	535		5.165			
MOTA	2222	OH2 WAT	536		35.064			
. ATOM	2223	OH2 WAT	537	:	7.785		•	1.00 19.77
MOTA	2224	OH2 WAT	538	•	2.503			1.00 23.38
MOTA	2225	OH2 WAT	539		2.763	-3.299	20.083	
MOTA	2226	OH2 WAT	54 0		6.475	6:912	39:440	1.00 22.13

FIG.11A-53

ATOM	2227	OH2 WAT	541	-6.228	9.593	24.818	1.00 26.15
MOTA	2228	OH2 WAT	542	37.153	5.154	30.029	1.00 23.86
MOTA	2229	OH2 WAT	54 3	8.552	2.510	13.829	1.00 21.71
ATOM	2230	OH2 WAT	544	16.101	3.059	45.670	1.00 22.52
MOTA	2231	OH2 WAT	545	32.130	14.940	31.845	1.00 22.82
MOTA	2232	OH2 WAT	546	18.050	14.095	15.782	1.00 22.03
ATOM	2233	OH2 WAT	547	24.287	11.877	41.531	1.00 25.65
MOTA	2234	OH2 WAT	:548	0.491	-4.750	31.613	1.00 21.18
MOTA	2235	OH2 WAT	549	7.787			1.00 23.30
MOTA	2236	OH2 WAT	550	12.435	-5.647	20.701	1.00 31.34
MOTA	2237	OH2 WAT	552	25.857			1.00 28.63
			553				1.00 36.96
MOTA				-4.014			1.00 24.61
MOTA				10.571		-16.930	1.00 26.27
MOTA	2241			14.828			
MOTA		· ·					1.00 27.12
MOTA		OH2 WAT					1.00 31.85
MOTA		OH2 WAT			and the second second		1.00 27.77
MOTA	2245	OH2 WAT					1.00 30.04
MOTA	2246						1.00 17.65
MOTA	2247						1.00 24.33
MOTA		OH2 WAT		* * *			1.00 24.76
MOTA		OH2 WAT	565	25.621			1.00 22.51
MOTA	2250				-1.136	43.949	1.00 22.61
MOTA		OH2 WAT					1.00 23.78
MOTA	2252		568			13.451	
MOTA		OH2 WAT	569				1.00 27.31
MOTA	2254				12.602	29.260	1.00 24.68
MOTA		OH2 WAT					1.00 24.44
MOTA				30.240			1.00 30.85
MOTA		OH2 WAT		3.021			1.00 33.27
ATOM	2258		574				1.00 34.06
ATOM	2259	•	575				1.00 24.07
MOTA	2260	•	576	19.000			1.00 32.41
MOTA	2261		577				1.00 31.82
MOTA	2262	•				41.174	
MOTA	2263		579	9.859		26.441	
MOTA	2264		580			14.768	
MOTA			581			12.412	
MOTA	2266		582			16.519	•
MOTA	2267		583				
MOTA	2268	OH2 WAT	584	17.449	-10.259	23.231	1.00 43.02

FIG.11A-54

MOTA	2269	OH2 WAT	585	-8.993	2.971	28.486	1.00 30.80
MOTA	2270	OH2 WAT	. 586	-0.139	-3.655	43.071	1.00 37.06
MOTA	2271	OH2 WAT	588	16.750	16.297	23.374	1.00 29.48
ATOM	2272	OH2 WAT	58 9	5.136	6.789	43.328	1.00 32.09
ATOM	2273	OH2 WAT	590	5.961	15.786	26.926	1.00 22.25
MOTA	2274	OH2 WAT	591	11.771	0.434	-14.604	1.00 25.03
MOTA	2275	OH2 WAT	592	20.674	-11.849	31.603	1.00 28.56
MOTA	2276	OH2 WAT	593	16.561	0.669	8.704	1.00 30.46
MOTA	2277	OH2 WAT	594	25.900	1.235	13.342	1.00 25.92
ATOM	2278	OH2 WAT	595	14.762	0.666	-11.939	1.00 27.07
MOTA	2279	OH2 WAT	596	19.928	0.579	42.222	1.00 33.09
ATOM	2280	OH2 WAT	597	2.749	-4.838	23.485	1.00 28.02
MOTA	2281	OH2 WAT	599	2.241	-12.981	17.063	1.00 32.27
ATOM	2282	OH2 WAT	600	17.311	-11.858	43.919	1.00 46.62
ATOM	2283	OH2 WAT	601	10.116	0.287	13.907	1.00 23.36
ATOM	2284	OH2 WAT	602	-5.766 ⁻	4.131	31.307	1.00 37.38
ATOM	2285	OH2 WAT	603	8.777	-6.752	16.659	1.00 36.23
MOTA	2286	OH2 WAT	604	2.780	13.085	33.578	1.00 56.17
ATOM	2287	OH2 WAT	605	13.505	-9.621	24.772	1.00 27.48
ATOM	2288	OH2 WAT	606	19.499	-8.171	22.784	1.00 35.47
ATOM	2289	OH2 WAT	607	18.981	6.434	6.609	1.00 39.96
ATOM	2290	OH2 WAT	609	19.617	1.498	-10.274	1.00 46.75
MOTA	2291	OH2 WAT	610	7.105	14.231	31.956	1.00 30.92
ATOM	2292	OH2 WAT	611	-2.597	7.596	24.441	1.00 52.94
MOTA	2293	OH2 WAT	612	38.962	0.347	34.269	1.00 28.21
MOTA	2294	OH2 WAT	613	34.567	6.357	38.002	1.00 53.59
ATOM	2295	OH2 WAT	614	19.967	5.584	-11.241	1.00 30.33
ATOM	2296	"OH2 WAT	615	0.984	14.444	27.279	1.00 41.31
ATOM	2297	OH2 WAT	616	31.944	18.357	34.770	1.00 56.14
ATOM	2298	OH2 WAT	617	23.842	3.527	43.838	1.00 39.09
ATOM	2299	OH2 WAT	618	24.265	-10.048	29.048	1.00 43.37
ATOM	2300	OH2 WAT	619	13.920	0.583	10.143	1.00 28.77
ATOM	2301	OH2 WAT	620	13.884	17.699	20.194	1.00 53.38
MOTA	2302	OH2 WAT	621	15.456	13.880	40.976	1.00 38.26
MOTA	2303	OH2 WAT	622	-4.209	10.695	27.546	1.00 31.65
ATOM	2304	OH2 WAT	623	9.422	15.446	37.303	1.00 35.02
MOTA	2305	OH2 WAT	624	28.277	9.830	16.219	1.00 32.83
MOTA	2306	OH2 WAT	625	-2.164	-0.376	23.957	1.00 32.44
MOTA	2307	OH2 WAT	626	13.795	-8.227	-22.617	1.00 47.18
MOTA	2308	OH2 WAT	627	12.663	-2.391	45.836	1.00 32.05
MOTA	2309	OH2 WAT	628	3.919	-11.060	32.966	1.00 44.73
MOTA	2310	OH2 WAT	629	-2.517	11.533	34.098	1.00 54.44

FIG.11A-55

		12	•			-	•
MOTA	2311	OH2 WAT	630	25.613	14.652	11.863	1.00 63.79
ATOM	2312	OH2 WAT	631	11.909	11.704	41.097	1.00 34.25
MOTA	2313	OH2 WAT	632	-1.360	10.995	26.456	1.00 38.16
MOTA	2314	OH2 WAT	633	31.933	5.045	17.791	1.00 39.91
ATOM:	2315	OH2 WAT	634	22.722	-5.823	24.321	1.00 28.24
ATOM	2316	OH2 WAT	635	16.867	10.054	41.617	1.00 32.20
MOTA	2317	OH2 WAT	636	-0.030	10.808	17.607	1.00 37.23
MOTA	2318	OH2 WAT	637	-2.623	-2.811	32.773	1.00 41.25
MOTA	2319	OH2 WAT	638	31.929	21.354	29.330	1.00 38.21
MOTA	2320	OH2 WAT	639	17.980	15.951	20.755	1.00 60.27
MOTA	2321	OH2 WAT	640	29.018	-3.356	20.263	1.00 36.21
MOTA	2322	OH2 WAT	641	20.664	16.288	14.235	1.00 42.55
MOŤA	2323	OH2 WAT	642	7.328	13.948	36.591	1.00 55.67
ATOM	2324	OH2 WAT	643	11.409	16.717	20.413	1.00 25.47
MOTA	2325	OH2 WAT	644	16.547	13.154	13.670	1.00 25.26
MOTA	2326	OH2 WAT	645	15.596	15.812	18.554	1.00 34.13
MOTA	2327	OH2 WAT	646	25.131	5.610	6.079	1.00 53.07
ATOM	2328	OH2 WAT	647	-3.556	15.275	34.402	1.00 61.62
MOTA	2329	OH2 WAT	648	10.229		19.982	
ATOM	2330	OH2 WAT	649	20.662	8.866	43.464	1.00 51.89
ATOM	2331	OH2 WAT	650	23.069		4	1.00 25.83
MOTA	2332	OH2 WAT	651	26.751		18.349	1.00 16.47
MOTA	2333	OH2 WAT	652	4.110	-8.428		
MOTA	2334	OH2 WAT	654	A CONTRACT OF STREET AND ADDRESS.	-14.479		1.00 33.14
ATOM	2335	OH2 WAT	655	13.831	16.895	27.725	1.00 39.59
ATOM	2336	OH2 WAT	656	13.478	5.441		1.00 41.26
MOTA	2337	OH2 WAT	657	14.527	-6.733	41.081	1.00 39.50
MOTA.	2338	OH2 WAT	658	12.344	-8.188	-4.840	1.00 31.36
MOTA	2339	OH2 WAT	659	2.335	0.119	-12.679	1.00 46.96
MOTA	2340	OH2 WAT	660	-4.072	8.903	35.840	1.00 33.73
MOTA	2341	OH2 WAT	661	11.199	-3.361	13.690	1.00 30.89
MOTA	2342	OH2 WAT	662	33.630	13.397	20.072	1.00 32.18
ATOM	2343	OH2 WAT	663	-8.225	5.595		1.00 42.51
ATOM	2344	OH2 WAT	664	4.851	• •		1.00 38.06
MOTA	2345	OH2 WAT	665		6.937	-3.912	1.00 45.24
MOTA	2346	OH2 WAT	666	16.913	-0.045	-10.717	1.00 42.04
MOTA	2347	OH2 WAT	667	29.488			1.00 47.43
MOTA	2348	OH2 WAT	668	23.202			1.00 36.53
MOTA	2349	OH2 WAT	669		-10.157		1.00 34.64
MOTA	2350	OH2 WAT	670	30.193	6.969		
MOTA	2351	OH2 WAT	671	9.581			1.00 38.51
MOTA	2352	OH2 WAT	672	3.957	11.310	37.024	
					_		

FIG.11A-56

				·				
ATOM	2353	OH2 WAT	673	23.314	-12.393	29.627	1.00 44.03	
MOTA	2354	OH2 WAT	674	29.567	-4.326	22.984	1.00 38.54	
MOTA	2355	OH2 WAT	675	20.341	-13.530	33.695	1.00 35,66	
MOTA	2356	OH2 WAT	676	24.115	-2.262	12.332	1.00 24.55	
MOTA	2357	OH2 WAT	677	21.496	16.243	18.532	1.00 38.18	
MOTA	2358	OH2 WAT	678	1.474	14.677	18.946	1.00 34.15	
MOTA	2359	OH2 WAT	679	22.623	10.998	43.542	1.00 34.98	
MOTA	2360	OH2 WAT	680	22.204 .	4.868	42.384	1.00 35.66	
MOTA	2361	OH2 WAT	681	4.974	18.238	22.943	1.00 43.25	
MOTA	2362	OH2 WAT	682	7.600	17.266	28.095	1.00 47.36	
MOTA	2363	OH2 WAT	683	9.887	-4.665	20.529	1.00 55.08	
ATOM	2364	OH2 WAT	684	34.174	16.468	30.910	1.00 59.36	
MOTA	2365	OH2 WAT	685	14.332	-9.413	41.717	1.00 44.97	
MOTA	2366	OH2 WAT	686	-6.650	-2.511	31.135	1.00 56.20	
MOTA	2367	OH2 WAT	687	3.069	14.962	28.974	1.00 53.45	
MOTA	2368	S S04	901	-0.036	-4.899	27.988	1.00 27.31	
MOTA	2369	01 S04	901	0.702	-5.486	26.855	1.00 27.32	
ATOM	2370	02 S04	901	0.883	-4.694	29.123	1.00 30.06	
MOTA	2371	O3 SO4	4 901	-1.115	-5.818	28.406	1.00 25.85	
MOTA	2372	04 504	4 .901	-0.628	-3.611	27.579	1.00 30.90	
END	•			(*				

FIG.11A-57

						to an an an exemption one and a	
ATOM	1	CB ALA	2	-1.758	8.559 -13.6	537 1.00 37.12	
ATOM	. 2	C ALA	. 2	0.707	8.098 -13.	· 	
ATOM	3	O ALA	2	0.588	7.652 -14.6		
ATOM	4	N ALA		-0.253	10.204 -12.		
MOTA	5	CA ALA		-0.489	8.748 -12.8		
ATOM	6	N VAL		1.848	8.047 -12.8		
MOTA	7	CA VAL		3.063	7.454 -13.3		
ATOM	8	CB VAL		4.313	7.955 -12.6	· ·	
MOTA	9	CG1 VAL		5.571	7.440 -13.3		
ATOM	10	CG2 VAL		4.317	9.475 -12.5		
ATOM	11	C VAL		2.978	5.903 -13.3		
ATOM	12	O VAL		2.931	5.330 -12.2		
ATOM	13	N PRO		2.991	5.224 -14.4		
ATOM	14	CD PRO	4	3.225	5.848 -15.7		
ATOM	15	CA PRO	4	2.907	3.767 -14.6		
ATOM	16	CB PRO	4	3.523	3.536 -15.9		
ATOM	17	CG PRO	4	2.992	4.691 -16.7		
ATOM	18	C PRO		3.439	2.787 -13.5		
ATOM	19	O PRO	4	2.692	1.913 -13.0		1
MOTA	20	N PHE		4.703	2.917 -13.1		214 214
ATOM	21	CA PHE	5	5.317	1.949 -12.2		
ATOM	22	CB PHE	5	6.565	1.362 -12.9		.:
ATOM	23	CG PHE	5	6.385	1.053 -14.3		i
ATOM	24	CD1 PHE	5	7.159	1.694 - 15.3		. •
ATOM	25	CD2 PHE	5	5.455	0.112 -14.8		
ATOM	26	CE1 PHE	5	7.001	1.390 -16.7		
MOTA	27	CE2 PHE		5.289	-0.198 -16.1		
MOTA	28	CZ PHE	5	6.067			
ATOM	29	C PHE	5	5.770	2.421 -10.8		
ATOM	30	O PHE		6.569	1.742 -10.2	26 1.00 29.38	
ATOM	31	N VAL		5.261	3.559 -10.4		
MOTA		CA VAL		5.665	4.110 -9.1		
ATOM	33	CB VAL	6	5.120		·	
MOTA	34	CG1 VAL	6	5.730	6.201 -7.7		
ATOM	35	CG2 VAL	6 -	5.439	6.368 -10.1		
MOTA	36	C VAL		5.270	3.291 -7.8		
MOTA	37	O VAL		5.731 ·			
MOTA	38	N GLU	7	4.441	2.268 -8.0		
ATOM	39	CA GLU	7	4.023 -			
ATOM -	40	CB GLU	7	2.536	1.131 -7.0		
ATOM	41	CG GLU	7 .	1.797	2.481 -6.8		

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MOTA	42	CD GLU	7.	2.340	3.275	-5.622	1.00 37.65
ATOM	43	OE1 GLU	7.	3.400		-5.755	1.00 38.88
ATOM	44	OE2 GLU	7.	1.728	3.259	-4.532	1.00 37.93
ATOM	45	C GLU	7	4.819	0.213	-6.723	1.00 29.89
ATOM	46	0 GLU	7	4.546	-0.563	-5.806	1.00 28.91
MOTA	47	N ASP	8	5.827	0.010	-7.566	1.00 27.92
MOTA	48	CA ASP	8	6.671	-1.163	-7.451	1.00 26.87
MOTA	49	CB ASP	8	7.122	-1.675	8.820	1.00 27.02
ATOM	50	CG ASP	8	5.988	-2.243	-9.636	1.00 28.55
ATOM	51	OD1 ASP	8	5.115	-2.957	-9.092	1.00 28.96
ATOM	52	OD2 ASP	- 8	5.984	-1.978	-10.856	1.00 29.04
MOTA	53	C ASP	8	7.902	-0.881	-6.651	1.00 26.17
MOTA	54	0 ASP	. 8-	8.599	0.112	-6.880	1.00 25.38
ATOM .	55	N TRP	9	8.165	-1.767	-5.698	1.00 25.50
ATOM	56	CA TRP	9	9.316	-1.674	-4.814	1.00 25.48
ATOM	57	CB TRP	9	8.856	-1.476	-3.360	1.00 26.35
ATOM	58	CG TRP	. 9 -	8.975	-0.060	-2.872	1.00 27.81
ATOM	59	CD2 TRP	9	7.939	0.920	-2.829	1.00 27.99
MOTA	60	CE2 TRP	9	8.511	2.110	-2.324	1.00 28.08
MOTA	61	CE3 TRP	9	6.580	0.910	-3.169	1.00 27.50
ATOM	62	CD1 TRP	9	10.108	0.557	-2.404	1.00 28.28
MOTA	63	NE1 TRP	9	9.837	1.860	-2.074	1.00 27.98
MOTA	64	CZ2 TRP	9	7.775	3.279	-2.150	1.00 27.32
MOTA	65	CZ3 TRP	9	5.848	2.070	-2.996	1.00 29.26
MOTA	66	CH2 TRP	9.	6.447	3.243	-2.488	1.00 29.79
MOTA	67	C TRP	9	10.129	-2.960	-4.877	1.00 25.40
ATOM	68	0 TRP	9	9.634	-4.028	-4.523	1.00 25.51
ATOM	69	N ASP	10	11.374	-2.857	-5.328	1.00 24.86
ATOM	70	CA - ASP-	10	12.260	-4.015	-5.414	1.00 25.35
MOTA	71	CB ASP	10	13.412	-3.734	-6.381	1.00 25.70
MOTA	72	CG ASP	10	12.908	-3.616	-7.816	
MOTA	7 3	OD1 ASP	10	13.473	-2.799	-8.588	
MOTA	. 74	OD2 ASP	10	11.959	-4.361	-8.157	1.00 26.29
ATOM	7 5	C ASP	10	12.875	-4.328		1.00 26.53
MOTA	. 76	0 ASP	10	13.358	-3.426		1.00 25.47
MOTA	77	N LEU	11	12.836		-3.645	
MOTA	78	CA LEU	11	13.415	-6.041	-2.373	1.00 29.13
ATOM	7 9	CB LEU	11	12.585	-7.191	-1.780	1.00 29.22
ATOM	80	CG LEU	11	11.165	-6.817	-1.329	1.00 29.89
ATOM	81	CD1 LEU	11	10.370	-6.260	-2.494	
ATOM	82	CD2 LEU	· 11	10.463	-8.052	-0.768	
ATOM	83	C. LEU	11	14.828	-6.461		1.00 30.25

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	MOTA	84	0	LEU	11	15.058	-7.557	-3.235	1.00 30.42
	MOTA	.85	N	VAL	. 12	15.783	-5.581	-2.436	1.00 30.42
	MOTA	86	CA	VAL	12	17.176	-5.813	-2.797	1.00 32.51
	MOTA	87	CB	VAL	12	17.798	-4.495	-3.312	1.00 33.09
	MOTA-	88	CG1	VAL	12	19.207	-4.740	-3.855	1.00 34.30
	MOTA	.89	CG2	VAL	12	16.907	-3.905	-4.400	1.00 33.06
	MOTA	90	С	VAL	12	18.130	-6.437	-1.774	1.00 33.49
	MOTA	91	0	VAL	12	19.202	-6.906	-2.151	1.00 33.55
	MOTA	92	N	GLN	13	17.767	-6.444	-0.495	1.00 34.17
	MOTA	93	CA	GLN	13	18.646		0.511	1.00 35.23
	MOTA	94	CB	GLN		19.962	-6.269	0.625	1.00 36.13
	MOTA	95	CG	GLN	13	19.957	•		1.00 37.34
	MOTA	96	CD	GLN	13	21.380	-4.268	0.950	1.00 38.25
•	ATOM	97	0E1	GLN	13	21.930	-4.495	2.029	1.00 38.73
	MOTA	98	NE2	GLN	13	21.981	-3.608	-0.039	1.00 39.12
	MOTA	99	C	GLN	13	18.064	-7.158	1.892	1.00 35.56
	MOTA	100	0	GLN	13	17.359	-6.266	2.362	1.00 35.04
	MOTA	101	. N	THR	14	18.359	-8.275	2.549	1.00 36.12
	MOTA	102		THR		17.871	-8.516	3.901	1.00 37.26
	MOTA	103	10.4	THR	14	18.035	-9.992	4.309	1.00 37.85
	ATOM	104		THR		17.689	-10.148	5.691	1.00 39.17
	MOTA	105		THR		19.471	-10.442	4.102	1.00 38.49
	ATOM	106		THR		18.653	-7.673	4.879	1.00 37.46
	MOTA	107		THR		19.864	-7.503	4.737	1.00 37.37
	MOTA	108		LEU	15	17.961	· · ·	5.872	1.00 37.76
	MOTA	109		LEU	15	18.604	-6.302	6.884	1.00 38.54
	MOTA	110		LEU		17.827	-5.003	7.100	
	MOTA	111		LEU		17.768		5.946	1.00 38.81
	ATOM	112	4.5	LEU		19.162		5.539	1.00 38.58
	ATOM	113		LEU	15	17.075	-4.674	4.787	1.00 39.25
	MOTA	114		LEU	15	18.662	-7.039	8.201	1.00 39.26
	ATOM	115		LEU		19.189		9.190	1.00 39.06
	ATOM	116		GLY		18.112	-8.248		1.00 40.02
	MOTA	117		GLY		18.101			1.00 41.67
	ATOM	118		GLY		16.767		9.569	1.00 42.84
	MOTA	119		GLY		15.726	-9.165	9.280	1.00 43.09
	MOTA	120		GLU	17		-11.003	10.024	1.00 43.99
	ATOM	121		GLU	17 ·		-11.775	10.203	1.00 44.96
	MOTA	122		GLU	17		-13.210	9.707	
	ATOM	123		GLU			-13.419	8.237	1.00 46.77
	ATOM	124		GLU	17		-14.915	8.042	1.00 47.36
	MOTA	125	UE1	GLU	17	17.089	-15.524	8.805	1.00 47.74

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MOTA	126	OE2 GLU	17	15.667 -15.486	7.134	1.00 48.12
MOTA	127	C GLU	17	15.131 -11.854	11.664	1.00 45.33
ATOM	128	0 GLU	17	15.952 -11.725	12.576	1.00 45.16
MOTA	129	N GLY	18	13.835 -12.066	11.872	1.00 45.83
MOTA	130	CA GLY	18	13.296 -12.170	13.216	1.00 46.41
MOTA	131	C GLY	18	12.627 -13.515	13.409	1.00 46.60
MOTA	132	0 GLY	18	12.340 -14.212	12.437	1.00 46.89
MOTA	133	N ALA	19	12.376 -13.880	14.663	1.00 46.88
ATOM	134	CA ALA	: 19	11.745 -15.155	14.977	1.00 46.85
MOTA	135	CB ALA	19	11.695 -15.350	16.492	1.00 46.77
MOTA"	136	C ALA	19	10.342 -15.268	14.390	1.00 46.78
ATOM	137	O ALA	19	9.701 -16.310	14.505	1.00 47.08
MOTA	138	N TYR	20	9.867 -14.196	13.763	1.00 46.62
MOTA	139	CA TYR	20	8.533 -14.187	13.165	1.00 46.18
MOTA	140	CB TYR	20	7.504 -13.682	14.185	1.00 46.74
MOTA	141	CG TYR	20	8.056 -12.671	15.169	1.00 47.27
MOTA	142	CD1 TYR	20	8.661 -11.493	14.728	1.00 47.21
MOTA	143	CE1 TYR	20	9.195 -10.576	15.629	1.00 47.56
MOTA	144	CD2 TYR	20	7.994 -12.904	16.544	1.00 47.53
MOTA	145	CE2 TYR	20	8.524 -11.993	17.454	1.00 47.75
MOTA	146	CZ TYR	20	9.125 -10.833	16.990	1.00 47.80
MOTA	147	OH TYR	20	9.673 -9.940	17.884	1.00 48.14
MOTA	148	C TYR	20	8.449 -13.336	11.886	1.00 45.48
MOTA	149	O TYR	20	7.509 -12.557	11.708	1.00 45.95
ATOM	150	N GLY	21	9.432 -13.496	11.004	1.00 44.27
MOTA	151	CA GLY	21	9.441 -12.742	9.761	1.00 42.92
MOTA	152	C GLY	21	10.817 -12.250	9.349	1.00 41.50
ATOM	153	O GLY	21	11.833 -12.774	9.800	1.00 41.92
MOTA	154	N GLU	22	10.850 -11.236	8.489	1.00 40.28
MOTA	155	CA GLU	22	12.111 -10.671	8.012	1.00 38.41
ATOM	156	CB GLU	22	12.571 -11.389	6.736	1.00 39.39
MOTA		CG GLU	22	11.451 -11.244	5.667	1.00 40.60
MOTA	158	CD GLU	22	11.755 -11.845	4.292	1.00 41.48
ATOM		OE1 GLU		10.825 -11.842	3.459	1.00 42.22
MOTA	160	OE2 GLU	22	12.882 -12.309	4.019	1.00 42.38
MOTA	161	C GLU	22	11.967 -9.174	7.684	1.00 36.25
ATOM	162	O GLU	22	10.858 -8. <i>6</i> 55	7.577	1.00 36.15
MOTA	163	N VAL	23	13.098 -8.497	7.537	1.00 34.42
ATOM	164	CA VAL	23	13.116 -7.077	7.198	1.00 32.22
ATOM	165	CB VAL	23	13.731 -6.227	8.338	1.00 32.25
ATOM	166	CG1 VAL	23	13.708 -4.749	7.965	1.00 30.96
ATOM	167	CG2 VAL	23	12.958 -6.456	9.622	1.00 31.12

ATOM	168	C VA	L 23	13.967	-6.926	5.945	1.00 31.31
MOTA	169	0 VA			-7.343	5.915	1.00 31.31
MOTA	170	N GL			-6.335	4.909	1.00 30.82
MOTA	171	CA GL			-6.148	3.647	1.00 30.37
MOTA	172	CB GL			-6.832	2.504	1.00 23.34
MOTA	173	CG GL			-8.363		1.00 33.39
ATOM	174	CD GL	N 24		-8.990	2.258	1.00 33.39
MOTA	175	OE1 GL	N - 24		-8.607		1.00 35.21
MOTA	176	NE2 GL	N 24	•	-9.958	3.101	1.00 35.38
MOTA	177	C GL	N 24	· ·		3.268	1.00 28.82
ATOM	178	O GL	N 24		-3.878	3.612	1.00 28.38
ATOM	179	N LE	U 25		-4.374		1.00 27.94
MOTA	180	CA LE	U 25	15.515	-3.015		1.00 27.18
ATOM .	181	CB LE	U 25	17.008	-2.729	•	1.00 28.17
MOTA	182	CG LE	U 25	17.457	-1.432	1.183	1.00 29.19
MOTA	183	CD1 LE	U 25	17.192	-1.511	-0.302	1.00 31.15
MOTA	184	CD2 LE		16.728	-0.244	1.790	1.00 29.21
MOTA		C LE		14.808	-2.978	0.765	1.00 26.32
MOTA	186			15.078	-3.808	-0.103	
MOTA	187	N AL		13.886	-2.037	0.608	1.00 25.18
MOTA	188	CA AL		13.134		-0.635	1.00 23.91
ATOM	189	CB AL		11.638	-1.992	-0.351	1.00 24.21
ATOM	190	C AL			-0.641	-1.335	1.00 23.39
ATOM	191	O AL	the second of the second	•	0.414	-0.713	1.00 23.17
MOTA	192	. N., YA			-0.719	-2.648	1.00 22.24
MOTA	193	CA VA			0.460	-3.433	
MOTA	194	CB VA			0.311	-4.031	1.00 21.43
MOTA	195	CG1 VA			1.514		1.00 21.22
MOTA	196	CG2 VA	the second secon		0.177		1.00 20.94
MOTA	197	C VA			0.020	-4.503	1.00 20.42
MOTA	٠.	O VA			-0.320	-5.223	1.00 20.31
MOTA	199	N AS			1.831	-4.606	1.00 19.21
MOTA	200	CA AS	•	•	2.108	-5.575	1.00 18.90
MOTA	201	CB AS	• .		3.511	-5.343	1.00 19.28
MOTA	202	CG AS			3.680	-6.151	1.00 19.66
MOTA	203	OD1 AS			4.128	-7.294	1.00 18.90
MOTA	204	ND2 AS			3.313		1.00 20.03
MOTA	205	C AS			1.962	-6.985	1.00 18.62
MOTA	206	O AS		12.893	2.482	-7.343	1.00 19.04
NOTA	207	N AR		_	1.253	-7.797	1.00 17.64
MOTA	208	CA AR			1.009	-9.185	1.00 17.42
MOTA	209	CB AR	G · 29	10.344	0.191	-9.860	1.00 17.14

ATOM	210	CG ARG	29	10.521	0.026 -11.359	1.00 16.26
MOTA	211	CD ARG	29	9.318	-0.788 -11.804	
MOTA	212	NE ARG	29	9.278	-0.897 -13.267	
MOTA	213	CZ ARG		8.410	-1.645 -13.940	
MOTA	214	NH1 ARG	29	7.484	-2.359 -13.289	
MOTA	215	NH2 ARG	29	8.481	-1.699 -15.267	
MOTA	216	C ARG	29	11.684	2.279 -9.978	
ATOM	217	0 ARG	29	12.641	2.364 - 10.759	
ATOM	218	N VAL	30	10.821	3.266 -9.769	
ATOM	219	CA VAL	30	10.901	4.540 -10.477	
ATOM	220	CB VAL		9.492	5.153 -10.639	
ATOM	221	CG1 VAL	30	9.587	6.584 -11.138	
ATOM	222	CG2-VAL		8.653		
ATOM	223	C VAL		11.797	5.605 -9.819	
ATOM	224	O VAL	30	12.723	6.110 -10.444	
ATOM	225	N THR		11.527	5.930 -8.555	
ATOM	226	CA THR	31	12.277	6.980 -7.868	
ATOM	227	CB THR		11.406	7.701 -6.820	
ATOM	228	OG1 THR	31	11.118	6.801 -5.742	
ATOM	229	CG2 THR	31	10.110	8.198 -7.448	
ATOM	230	C THR	31	13.559	6.616 -7.144	
ATOM	231	O THR	31	14.339	7.507 -6.784	
ATOM	232	N GLU	32	13.767	5.319 -6.934	
ATOM	233	CA GLU	32	14.927	4.772 -6.238	
ATOM	234	CB GLU	32	16.239	5.257 -6.861	
ATOM .	235	CG GLU	- 32	16.258	4.557 -8.256	
MOTA	236	CD GLU	32	17.606	4.686 -8.924	
ATOM	237	OE1 GLU	32	18.624	4.258 -8.330	
ATOM	238	OE2 GLU	. 32	17.636	5.212 -10.049	
MOTA	239	C GLU	32	14.912	5.113 -4.755	
ATOM	240	O GLU	32	15.918		
ATOM	241	N GLU	. 33	13.770	5.584 -4.271	
ATOM	242	CA GLU	• 33	13.632	5.907 -2.851	
ATOM	243	CB GLU	33	12.281	6.581 -2.581	•
ATOM	244	CG GLU	33 ·	11.945	6.632 -1.055	
ATOM	245	CD GLU	33	10.594	7.270 -0.711	
ATOM	246	OE1 GLU	33 .	9.689	7.294 -1.571	•
MOTA	247	OE2 GLU	33	10.425	7.742 0.437	
MOTA	248	C GLU	33	13.727	4.575 -2.097	,
MOTA	249	O GLU	33	13.229	3.546 -2.575	_ , , ,
ATOM .	250	N ALA	34 [.]	14.373	4.595 -0.931	•
ATOM	251	CA ALA	34	14.537	3.388 -0.132	

ATOM	252	· CB	ALA	34	16.006	3.207	0.244	1.00 25.80
ATOM	253	C	ALA	34	13.694	3.404	1.125	1.00 25.80
ATOM	254	.0	ALA	34	13.559	4.435	1.785	1.00 25.49
ATOM	255	N	VAL	35	13.117	2.254	1.449	1.00 25.71
ATOM	256	CA	VAL	35	12.324	2.110	2.666	1.00 25.20
ATOM	257	CB	VAL	35	10.799	2.237	2.414	1.00 25.48
ATOM	258	CG1		35	10.461	3.621	1.870	1.00 25.96
MOTA	259	CG2	VAL	35	10.339	1.150	1.467	1.00 27.05
ATOM	260	·C	VAL	35	12.585	0.738		1.00 25.10
MOTA	261	0	VAL	35 4	13.041	-0.129	2.433	1.00 25.54
ATOM	262	N	ALA	36	12.324		4.465	1.00 24.18
MOTA	263	CA	ALA	.36	12.503	-0.793		1.00 23.82
MOTA	264	CB	ALA	36	13.081	-0.671		1.00 23.63
ATOM	265	C	ALA	36	11.143	-1.443		
ATOM	266	0	ALA	36	10.147	-0.807		1.00 23.23
ATOM	267	N	VAL	37	11.087	-2.714	4.719	1.00 24.59
ATOM	268	CA	VAL	37	9.832	-3.439	4.718	1.00 25.64
ATOM	269	CB	VAL	37 🚕	9.506	-4.006	3.316	1.00 25.94
MOTA	270		VAL	37	8.174	-4.755		
MOTA	271	CG2	VAL	37	9.453	-2.870	2.311	1.00 26.82
MOTA	272	*,-	VAL	37	9.915	-4.556	5.683	1.00 26.26
MOTA	273	0	VAL	37	10.724	-5.462	5.532	1.00 25.86
ATOM	274		LYS	38	9.091	-4.489	6.716	1.00 27.45
ATOM	275	CA	LYS	38	9.066	-5.534	7.725	1.00 29.32
ATOM	276	CB	LYS	38	8.729	-4.923	9.091	1.00 29.33
ATOM	277	CG	LYS	38	8.764	-5.913		1.00 30.35
ATOM	278		LYS	38	8.546	-5.119	11.550	1.00 30.39
ATOM	279	CE	LYS	38	8.715		12.796	1.00 30.62
ATOM	280		LYS	38	8.705	-5.183	14.038	1.00 29.08
MOTA	281		LYS	38	8.007	-6.537	7.295	1.00 30.44
ATOM	282	٠.	LYS	38	6.824	-6.212	7.247	1.00 30.48
ATOM		N	ILE	39	8.443	-7.748	6.964	1.00 32.39
ATOM	284		ILE	39	7.539	-8.805		1.00 34.05
MOTA	285	CB	ILE	39	8.130	-9.560		
MOTA	286		ILE	39		-10.462	4.702	1.00 34.87
MOTA	287		ILE		8.603	-8.553	4.271	1.00 34.64
MOTA	288		ILE	39		-9.188	3.005	1.00 35.49
MOTA	289	C	ILE	39	7.295	-9.775	7.672	1.00 35.39
MOTA	290	0	ILE	39		-10.291	8.273	1.00 35.78
MOTA	291	N	VAL	· 40	_	-10.018	7.977	1.00 36.68
MOTA	292	CA	VAL	40		-10.919	9.069	1.00 38.18
MOTA	293	CB	VAL	40	5.304	-10.116	10.337	1.00 38.57

FIG.11B-7

ATOM	294	CCI	VAL	40	£ 530 0 000		
ATOM	295		VAL	40	6.530 -9.382		1.00 38.14
ATOM	296	C	VAL	40	4.204 -9.118		1.00 38.65
ATOM	297	0	VAL		4.514 -11.850	8.710	1.00 39.27
ATOM	298	N	ASP	40	3.393 -11.401	8.470	1.00 39.49
ATOM	299	CA	ASP	41	4.794 -13.150	8.678	1.00 40.75
ATOM				41	3.772 -14.145	8.355	1.00 42.21
	300	CB	ASP	41	4.412 -15.507		1.00 42.73
MOTA	301	CG	ASP	41	3.334 - 16.461	7.562	1.00 43.32
MOTA	302		ASP	41	2.325 -16.676	8.268	1.00 44.00
ATOM	303		ASP	41	3.499 -16.997	6.445	1.00 43.83
MOTA	304	C	ASP	41	2.790 -14.290	9.515	1.00 42.79
ATOM	305	0	ASP.	41	3.155 -14.759	10.596	1.00 43.26
ATOM	306	N	MET	42	1.542 -13.895	9.278	1.00 43. 17
ATOM	307	CA	MET	42	0.495 -13.954	10.295	1.00 43.48
ATOM	308	CB	MET	42	-0.859 -13.567	9.690	1.00 44.53
MOTA	309	CG	MET	42	-0.894 -12.152	9.054	1.00 45.50
ATOM	310	SD	MET	4 2	-2.480 -11.758	8.271	1.00 47.53
ATOM	311	CE	MET	42	-3.298 -10.869	9.602	1.00 45.69
MOTA	312	C	MET	42	0.324 -15.322	10.981	1.00 43.53
ATOM	313	0	MET	42	-0.488 -15.458	11.898	1.00 43.38
MOTA	314	N	ALA	43	1.087 -16.320	10.543	1.00 43.07
ATOM	315	CA	ALA	43	0.991 -17.656	11.125	1.00 43.42
MOTA	316	CB	ALA	43	-0.040 -18.484	10.359	1.00 43.10
ATOM	317	:-C	ALA	43	2.327 -18.378	11.137	1.00 43.21
ATOM	318	0	ALA	43	2.386 -19.594		1.00 43.78
MOTA	319	N	ALA	44	3.403 -17.633	11.357	1.00 42.71
ATOM	320	CA	ALA	44	4.733 -18.227	11.388	1.00 42.36
ATOM	321	CB	ALA	44	5.750 -17.260		1.00 41.81
ATOM	322	C	ALA	44	5.117 -18.581	12.817	1.00 42.44
MOTA	323	0	ALA	44	6.254 -18.285	13.244	1.00 42.57
ATOM	324	OT	ALA	44	4.263 -19.178	13.504	1.00 42.66
ATOM	325	CB	CYS	48	0.655 -13.396	16.575	1.00 44.68
MOTA	326	SG	CYS	48	-0.887 -13.047	17.451	1.00 46.03
ATOM	327	C	CYS	48	-0.464 -12.204	14.719	1.00 43.08
MOTA	328	0	CYS.	48	0.015 -11.080	14.888	1.00 42.59
ATOM	329	N	CYS	48	1.582 -13.589	14.278	1.00 42.75
ATOM	330	CA	CYS	48		15.083	1.00 42.75
ATOM	331	N .	PRO .	. 49	-1.700 -12.387	14.224	1.00 43.49
ATOM	332	CD	PRO	. 49	-2.344 -13.692	13.989	
ATOM	333		PRO		-2.591 -11.294		1.00 42.03
ATOM	334	CB	PRO:	49	-3.891 -12.028	13.824	1.00 42.17
ATOM	335	CG	PRO ·	49		13.488	1.00 42.12
21.011	555	Ju	. 110	77	-3.406 -13.342	12.979	1.00 41.95

MOTA	336	C PRO	4 9	-2.745	-10.224	14.887	1.00 41.81
MOTA	337	O PRO	49	-2.664	-9.031		1.00 42.45
MOTA	338	N GLU	50	-2.969	-10.652	16.126	1.00 41.02
MOTA	339	CA GLU	50	-3.156	-9.729	17.246	1.00 39.67
MOTA	340	CB GLU	50	-3.526	-10.488	18.522	1.00 40.23
MOTA	341	CG GLU	50	-3.682	-9.495	19.715	1.00 40.60
MOTA	342	CD GLU	50	-3.534	-10.215		1.00 41.12
MOTA	343	OE1 GLU	50		-10.793	21.314	1.00 42.36
MOTA	344	OE2 GLU	50	•	-10.196	21.839	1.00 42.30
MOTA	345	C GLU	50	-1.926			1.00 38.68
MOTA	346	O GLU	50	-2.025	-7.666	17.747	1.00 38.74
ATOM	347	N ALA	51	-0.775		17.714	1.00 37.63
ATOM	348	CA ALA	51		-8.856	18.055	1.00 36.16
ATOM	349	CB ALA	51	1.591	-9.872		1.00 36.72
ATOM	350	C ALA	51	, .		17.010	1.00 35.47
MOTA	351	O ALA	51	1.150		17.330	
MOTA	352			0.885		15.754	
MOTA	353	CA ILE	52		-7.370		1.00 33.77
MOTA	354	CB ILE	52	1.420	-8.194	13.354	1.00 34.35
MOTA	355	CG2 ILE	52	0.066	-8.432	1	1.00 34.21
ATOM	356	CG1 ILE	52	2.451	-7.522		
ATOM	357	CD1 ILE	52		-6.082	12.050	
MOTA	358	C ILE	52		-6.248		
ATOM	359	0 ILE	52	0.462	-5.095	14.214	1.00 32.57
ATOM	360	N LYS	53	-1.092	-6.600	14.826	1.00 32.29
MOTA		CA LYS	53	-2.177	-5.627	14.744	1.00 31.31
MOTA	362	CB LYS	53	-3.520	-6.275	15.097	1.00 32.79
MOTA	363	CG LYS	53	-4.591	-5.745	14.135	1.00 35.30
ATOM	364	CD LYS	53	-4.333	-6.330	12.710	1.00 36.83
ATOM	365	CE LYS	53	-5.147	-5.687	11.568	1.00 37.66
ATOM	366	NZ LYS	53	-4.748	-4.262	11.361	1.00 39.01
ATOM	367	C LYS	5 3	-1.922	-4.469	15.733	1.00 29.81
MOTA	368	O LYS	53	-2.123	-3.297	15.410	1.00 29.51
MOTA	369	N LYS	54	-1.471	-4.810	16.933	1.00 28.14
MOTA	370	CA LYS	54	-1.202	-3.801	17.942	1.00 26.62
ATOM	371	CB LYS	- 54	-0.984	-4.475	19.292	1.00 26.85
ATOM	372	CG LYS	54	-0.815	-3.468	20.426	1.00 26.40
MOTA	373	CD LYS	54	-0.807	-4.242	21.744	1.00 27.28
MOTA	374	CE LYS	54	-0.732	-3.338	22.970	1.00 27.15
ATOM	375	NZ LYS	54	-0.636	-4.143	24.224	1.00 27.93
MOTA	376	C LYS	54	0.008	-2.953	17.542	1.00 25.67
MOTA	377	0 LYS	54	0.027	-1.738	17.751	1.00 25.37

FIG.11B-9

ATOM	378	N G	LU 55		1.010	-3.599	16.950	1.00 24.57
ATOM	379	CA G	LU 55		2.210	-2.892	16.517	1.00 24.57
MOTA	380	CB G	LU 55		3.246	-3.892	15.987	1.00 22.70
ATOM	381	CG G	LU 55		4.551	-3.198	15.516	1.00 22.01
ATOM	·382	CD G	LU 55		5.645	-4.208	15.112	1.00 22.01
ATOM	383	OE1 G			5.523	-5.412	15.423	1.00 22.15
MOTA	384	OE2 GI			6.643	-3.798	14.487	1.00 23.29
ATOM	385		U 55		1.842	-1.857		1.00 22.01
MOTA	386		JU 55		2.387	-0.756	15.399	1.00 22.77
MOTA	387		E 56		0.898	-2.215	14.570	1.00 22.54
MOTA	388		E 56		0.467	-1.320	13.507	1.00 22.45
MOTA	389	CB II	E 56		-0.378	-2.105	12.475	1.00 23.28
ATOM-	390	CG2 II		•	-0.995	-1.170	11.459	1.00 23.28
MOTA	391	CG1 II		131	0.516	-3.127	11.778	1.00 22.95
ATOM	392	CD1 II	E 56		-0.237	-4.041		
MOTA	393	C II	E 56		-0.299		14.087	1.00 22.53
MOTA	394	0 II	E 56		-0.092	0.979	13.712	1.00 22.30
MOTA	395	N C	/S 57		-1.179	-0.493	15.030	1.00 22.44
ATOM	396	CA CY	/S 57		-2.008	0.497	15.709	1.00 23.00
MOTA	397	CB CY	S 57		-2.832	-0.188	16.804	1.00 23.58
MOTA	398	SG CY	/S · 57		-3.925	0.986	17.618	1.00 26.04
ATOM	399	C C	(S 57		-1.157	1.603	16.347	1.00 22.83
MOTA	400	O CY	S 57		-1.441	2.795	16.203	1.00 23.71
MOTA	401	N II	E 58		-0.115	1.187		1.00 21.58
MOTA	402	CA II	E 58		0.757	2.129		1.00 20.88
MOTA	403	CB II	E 58		1.594	1.391	18.786	1.00 21.07
MOTA	404	CG2 IL	E 58		2.703	2.311	19.326	1.00 20.39
MOTA	405	CG1 II	E 58		0.661	0.937	19.916	1.00 21.01
ATOM	406	CD1 IL	E 58	<u>_:</u>	1.368	0.216	21.094	1.00 21.40
MOTA	407	C IL	E 58		1.583	2.907	16.737	1.00 19.99
MOTA	408	O II	E 58	·	1.747		16.888	
ATOM	409	N AS	N 59	٠.,	2.092	2.244	15.706	
MOTA	410		N 59		2.883	2.955		1.00 20.70
ATOM	411		N 59		3.358			1.00 19.66
MOTA	412	CG AS	N 59		4.803	1.554		1.00 19.76
ATOM	413	OD1 AS		,	5.736	2.319	13.609	1.00 21.26
ATOM	414	ND2 AS	N 59		4.985	0.321	14.287	1.00 19.87
ATOM	415	C AS	_		2.045	4.083	14.048	1.00 21.81
ATOM	416	0 · AS			2.567	5.147	13.720	1.00 21.63
MOTA	417	N · LY	S 60		0.752	3.836	13.864	1.00 22.44
MOTA	418	CA LY			-0.118	4.839	13.249	1.00 23.91
ATOM	419	CB LY	S 60		-1.528	4.280	13.027	1.00 24.44

FIG.11B-10

MOTA	420	CG LYS	60	-1.552	3.237	11.885	1.00 27.44
ATOM	421	CD LYS		-2.997	2.663	11.665	1.00 29.48
ATOM	422	CE LYS		-4.024	3.744	11.233	1.00 30.95
MOTA	423	NZ LYS		-5.377	3.169	10.933	1.00 32.88
MOTA	424	C LYS		-0.251	6.101	14.078	1.00 24.30
ATOM	425	O LYS		-0.657		13.574	1.00 24.07
MOTA	426	N MET		0.104		15.354	1.00 24.01
MOTA	427	CA MET		-0.002	· ——·	16.244	1.00 25.12
MOTA	428	CB MET	61	-0.249	6.693	17.676	1.00 25.40
MOTA	429	CG MET	61	-1.470	5.835	17.988	1.00 26.45
MOTA	430	SD MET	61	-1.392	5.217	19.669	1.00 29.36
MOTA	431	CE MET	61	-1.535	6.797	20.599	1.00 28.93
MOTA	432	C MET	61	1.255	8.008	16.297	1.00.24.94
MOTA	433	O MET	61	1.218	9.153	16.749	
ATOM	434	N LEL	62	2.359	7.458	15.809	1.00 24.38
MOTA	435	CA LEU	62	3.651	8.133	15.886	1.00 24.10
MOTA		CB LEL	62	4.742	7.099	16.141	1.00 24.29
ATOM		CG LEL	62	4.251	6.128	17.219	1.00 24.77
MOTA		CD1 LEL		5.273	5.004	17.283	1.00 24.61
ATOM		CD2 LEU		4.088	6.800	18.578	1.00 24.56
ATOM	of the factor of the first	C LEL	and the second of the second o	4.141	8.977	14.723	1.00 23.68
MOTA	4.1	0 LEU		3.965	8.615	13.556	1.00 23.81
MOTA		n asn		4.783		15.062	1.00 22.77
ATOM		CA ASN	general de la companya de la company		11.007	14.084	1.00 22.30
ATOM		CB ASN		4.255	11.825	13.407	1.00 24.66
MOTA		CG ASN			12.770		1.00 26.67
MOTA		OD1 ASN		•	and the second second	11.719	
MOTA		ND2 ASN	en e la me r de l'Agente	4.140		12.177	1.00 28.45
MOTA		C ASN		6.385		14.801	
MOTA		O ASN		6.037		15.363	1.00 19.93
MOTA		N HIS		7.645		14.795	1.00 18.13
MOTA		CA HIS		8.696	•	15.459	1.00 17.11
ATOM		CB HIS		8.666	11.908		1.00 16.62
MOTA		CG HIS		9.600	12.744	17.769	1.00 15.90
MOTA		CD2 HIS		9.402		18.439	1.00 15.85
MOTA		ND1 HIS		10.934	12.438	17.917	1.00 16.76
MOTA	•	CE1 HIS		11.519	13.373	18.642	1.00 16.09
MOTA		NE2 HIS		10.611	14.275	18.971	1.00 15.69
MOTA.		C HIS		10.038	11.910	14.827	1.00 17.07
MOTA	459			10.278	10.781	14.397	1.00 16.68
MOTA	460 461			10.918	12.908	14.771	1.00 16.86
MOTA	461	CA GLU	65	12.227	12.746	14.142	1.00 17.23

FIG.11B-11

MOTA	462	CB	GLU	65	12.977	14.081	14.097	1.00 19.17
MOTA	463	CG	GLU	65	13.115	14.589	15.530	1.00 23.09
MOTA	464	CD	GLU	65	12.010	15.573	15.921	1.00 24.78
MOTA	465	0E1	GLU	65	10.804	15.418	15.625	1.00 26.63
MOTA	466	0E2	GLU	65	12.412	16.555	16.575	1.00 29.13
MOTA	467	C	GLU	65	13.165	11.705	14.764	1.00 16.61
MOTA	468	0	GLU	65	14.136	11.290	14.123	1.00 15.95
ATOM	469	N	ASN	66	12.881	11.287	15.999	1.00 15.86
ATOM	470	CA	ASN	66	13.718	10.276	16.645	1.00 15.10
MOTA	471	CB	ASN	66	14.251	10.768	17.999	1.00 14.82
ATOM	472	CG	ASN	66	15.223	11.978	17.803	1.00 14.88
MOTA	473	OD1	ASN	66	14.921	13.102	18.214	1.00 15.31
MOTA	474	ND2	ASN	66	16.373	11.732	17.171	1.00 14.43
MOTA	475	C	ASN	66	12.968	8.975	16.839	1.00 15.46
ATOM	476	0	ASN	66	13.285	8.192	17.740	1.00 14.42
ATOM -	477	N	VAL	67	11.976	8.742	15.980	1.00 15.06
MOTA	478	CA	VAL	67	11.188	7.519	16.015	1.00 14.66
ATOM	479	CB	VAL	67	9.752	7.773	16.576	1.00 14.46
MOTA	480	CG1	VAL	67	8.896	6.527	16.418	1.00 14.49
ATOM	481	CG2	VAL	67	9.817	8.155	18.064	1.00 14.92
ATOM	482	C	VAL	67	11.079	7.032	14.567	1.00 15.45
ATOM	483	0	VAL	67 .	10.730	7.812	13.682	1.00 15.32
MOTA	484	N	VAL	68	11.398	5.762	14.326	1.00 14.74
ATOM	485	CA	VAL	68	11.318	5.209	12.968	
ATOM	486	CB	VAL	68	11.621	3.688	12.985	1.00 14.98
ATOM	487	CG1	VAL	68	11.344	3.072	11.604	1.00 15.35
MOTA	488	CG2	VAL	68	13.087	3.466	13.331	1.00 13.74
ATOM	489	C	VAL	68 .	9.953	5.508	12.374	1.00 16.26
ATOM	490	0 -	"VAL"	68	8.932	5.061	12.890	1.00 16.70
MOTA	491	N	LYS	69	9.939	6.255	11.272	1.00 17.50
ATOM	492	CA	LYS	69	8.688	6.638	10.629	
ATOM	493	CB	LYS	69	8.948		9.496	1.00 22.64
ATOM	494	CG.	LYS	69	9.172	9.162	9.649	
ATOM	495	CD	LYS	:69	10.454	9.843	10.196	
MOTA	496	CE	LYS	69	10.263	11.379	10.284	1.00 29.24
ATOM	497	NZ	LYS	69	11.485	12.071	10.783	1.00 31.62
ATOM .	498	C	LYS	69	7.927	5.460	10.056	1.00 20.86
ATOM	499	0	LYS	69	8.526	4.540	9.506	1.00 19.95
MOTA	500	N	PHE	· 70	6.605	5.497	10.204	1.00 21.61
MOTA	501	CA	PHE	70	5.708	4.465	9.696	1.00 22.90
MOTA	502	CB	PHE	70	4.624	4.154	10.731	1.00 23.86
ATOM	503	CG	PHE	70	3.610	3.142		1.00 25.07

					•				
	ATOM	504	CD1 PHE	70	3.984	1.828	10.020	1.00 25.78	
	MOTA	505	CD2 PHE	70	2.272	3.496	10.149	1.00 25.74	
	MOTA	506	CE1 PHE	70	3.038	0.873	9.652	1.00 27.17	
	ATOM	507	CE2 PHE	70	1.310	2.547	9.780	1.00 26.29	
	MOTA	508	CZ PHE	70	1.695	1.237	9.534	1.00 26.19	
	MOTA	509	C PHE	70	5.020	5.030		1.00 23.33	
	MOTA	510	O PHE	70	4.312	6.038	_	1.00 22.78	
٠.	ATOM	511	N TYR	71	5.226			1.00 24.38	
	MOTA	512	CA TYR	71	4.639	4.836	6.037	1.00 26.22	
	MOTA	513	CB TYR	71	5.615	4.622	4.886	1.00 25.69	
	ATOM	514	CG TYR	71	6.947	5.313		1.00 25.34	
	ATOM	515	CD1 TYR	71	7.023	6.687	-		
	MOTA	516	CE1 TYR	- 71	8.263	7.337		1.00 25.78	
	ATOM	517	CD2 TYR	71	8.139	4.606		1.00 25.28	
	ATOM	518	CE2 TYR	71	9.372	5.243		1.00 25.14	
	ATOM	519	CZ TYR	71	9.427	6.608		1.00 25.13	
	MOTA	520	OH TYR	71	10.653			1.00 26.09	٠
	MOTA	521	C TYR		3.327			1.00 27.64	
	MOTA	522	O TYR	71	2.579	4.675		1.00 28.91	
	ATOM	523	N GLY	72	3.044	3.016	and the second s	1.00 28.78	
	ATOM	524	CA GLY	72	1.814	2.306		1.00 30.86	
	MOTA	525	C GLY	72	1.968	0.802		1.00 31.92	
	ATOM	526	O GLY	72	3.057	0.297		1.00 31.91	
	MOTA	527	N HIS	73	0.872	0.080	5.862	1.00 33.79	
	ATOM	528	CA HIS	73	0.900	-1.376		1.00 35.69	
	ATOM	529	CB HIS	73	0.508	-1.844		1.00 35.93	
	ATOM	530	CG HIS	73	-0.894	-1.487			
	ATOM	531	CD2 HIS	73	-1.460	-0.295			
	ATOM	532	ND1 HIS	73	-1.900	-2.424		1.00 35.93	
	MOTA	533	CE1 HIS	73	-3.025	-1.825	8.163	1.00 36.12	
	MOTA	534	NE2 HIS	73	-2.785	-0.533		1.00 36.36	
	ATOM	5 35	C HIS	73	-0.058	-1.992	4.924	1.00 37.26	
	ATOM	536 :	O HIS	73	-1.020	-1.351	4.503	1.00 37.54	
	ATOM	537	N ARG	74	0.215	-3.236	4.542	1.00 39.37	
	ATOM	538	CA ARG	74	-0.617	-3.957	3.582	1.00 41.57	
	ATOM	539	CB ARG	74	0.193	-4.335	2.342	1.00 42.35	
	MOTA	540	CG ARG	74	0.662	-3.169		1.00 43.17	
	MOTA	541	CD ARG.	74	1.492	-3:729	0.290	1.00 43.98	
	ATOM	542	NE ARG	74	0.755	-4.738	-0.469	1.00 44.77	
	MOTA	· 543	CZ ARG	74	1.259	-5.412	-1.497	1.00 44.92	
	MOTA	544	NH1 ARG	74	2.505	-5.184	-1.892	1.00 45.32	
	MOTA	545	NH2 ARG	74	0.523	-6.320	-2.126	1.00 45.08	

MOTA	546	C	ARG	74	1.194	-5.241	4.175	1.00 42.67
MOTA	547	0	ARG	74	-0.714	-5.738	5.196	1.00 43.41
MOTA	548	N	ARG	75	-2.219	-5.773	3.513	1.00 43.71
MOTA	549	CA	ARG	75	-2.904	-6.993	3.945	1.00 44.56
MOTA	550	CB	ARG	75	-4.356	-6.992	3.455	1.00 44.64
MOTA	551	CG	ARG	75	-5.231	-5.765	3.791	1.00 45.27
MOTA	552	CD	ARG	75	-5.688	-5.646	5.256	1.00 45.40
MOTA	553	NE	ARG	75	-6.632	-6.700	5.620	1.00 45.56
MOTA	554	CZ	ARG	75	-7.160	-6.844	6.831	1.00 45.42
MOTA	555	NH1	ARG	75	-6.835	-6.002	7.804	1.00 45.59
MOTA	556	NH2	ARG	75 .	-8.021	-7.825	7.071	1.00 45.46
MOTA	557	C	ARG	75	-2.253	-8.266	3.377	1.00 44.76
MOTA	558	0	ARG	75	-1.782	-9.124	4.124	1.00 45.26
MOTA	559	N	GLU -	76	-2.247	-8.370	2.052	1.00 44.88
ATOM	560	CA	GLU.	76	-1.680	-9.517	1.346	1.00 44.86
ATOM	561	CB	GLU	76	-0.152	-9.439	1.337	1.00 45.21
MOTA	562	CG	GLU	76	0.450	-10.469	0.334	1.00 45.31
MOTA	563	CD	GLU	76	0.050	-10.137	-1.107	1.00 45.52
ATOM	564	0E1	GLU	76	0.511	-9.104	-1.639	1.00 45.49
ATOM	565	0E2	GLU	76	-0.731	-10.912	-1.704	1.00 45.23
ATOM	566	C	GLU	76	-2.116	-10.859	1.960	1.00 44.61
ATOM	567	0	GLU	76	-1.297	-11.758	2.171	1.00 44.52
ATOM	568	N	GLY	. 77	-3.409	-10.977	2.247	1.00 44.26
ATOM	569	CA	GLY	77 .	-3.938	-12.204	2.819	1.00 43.80
ATOM	570	. C	GLY	77	-3.414	-12.551	4.202	1.00 43.49
ATOM	571	0	GLY	77	-3.923	-12.054	5.208	1.00 43.67
ATOM	572	N	ASN.	78		-13.415	4.251	1.00 43.27
ATOM	573	CA	ASN	78 .		-13.841	5.517	1.00 43.15
ATOM	574	CB	ASN	78	-1.743	-15.371	5.593	1.00 43.84
ATOM	575	CG	ASN	78		-15.993	5.490	1.00 44.43
ATOM	576	0D1	ASN	78	-3.797	-15.925	4.444	1.00 44.43
MOTA	577	ND2	ASN	78	-3.608	-16.593	6.583°°	1.00 44.71
MOTA	578	C	ASN	78	-0.382	-13.285	5.753	1.00 42.53
MOTA	579	0	ASN	78	0.299	-13.684	6.699	1.00 43.03
MOTA	580	N.	ILE	79	0.061	-12.378	4.888	1.00 41.52
MOTA	581	CA	ILE	79	1.387	-11.782	5.035	1.00 40.34
ATOM	582	CB	ILE	79	2.264	-12.027	3.778	1.00 40.55
MOTA	583 ⁻	CG2	ILE	79	3.645	-11.411	3.973	1.00 40.53
MOTA	584		ILE	79	2.415	-13.530	3.519	1.00 40.50
ATOM.	585	CD1	ILE	79	3.142	-14.304	4.645	1.00 40.17
MOTA	586	C	ILE	79	1.243	-10.281	5.242	1.00 39.50
MOTA	587	0	ILE	79	0.747	-9.578	4.362	1.00 39.47

ATOM	588	N GLN	80	1.658	-9.794	6.411	1.00 38.59
ATOM	589	CA GLN	. 80	1.584	-8.368	6.729	1.00 37.48
MOTA	590	CB GLN	80	1.413	-8.135	8.237	1.00 38.53
MOTA	591	CG GLN	80	0.116	-8.469	8.997	1.00 39.05
MOTA	592	CD GLN	80	-1.025	-7.622	8.446	1.00 39.69
ATOM	593	OE1 GLN	80	-0.923	-6.397	8.363	1.00 40.14
ATOM	594	NE2 GLN	80	-2.119	-8.277	8.069	1.00 39.68
MOTA	595	C GLN	80	2.865	-7.668	6.312	1.00 36.30
MOTA	596	O GLN	80	3.963	-8.158	6.570	1.00 36.40
MOTA	597	N TYR	81	2.721	-6.518	5.662	1.00 34.84
MOTA	598	CA TYR	81	3.870	-5.741	5.194	1.00 33.41
MOTA	599	CB TYR	81	3.809	-5.546	3.674	1.00 33.77
MOTA	600	CG TYR	81	4.000	-6.809	2.856.	1.00 34.37
MOTA	601	CD1 TYR	81	5.272	-7.243	2.492	1.00 34.27
MOTA	602	CE1 TYR	81	5.452	-8.406	1.740	1.00 34.76
ATOM	603	CD2 TYR	81	2.903	-7.571	2.450	1.00 34.73
ATOM	604	CE2 TYR	81	3.074	-8.735	1.701	1.00 34.90
ATOM	605	CZ TYR	81	4.349	-9.145	1.350	1.00 34.65
MOTA	606	OH TYR	81	4.517	-10.296	0.614	1.00 35.49
ATOM	607	C TYR	81	3.873	-4.343	5.834	1.00 31.88
MOTA	608	O TYR	81	2.965	-3.548	5.602	1.00 32.17
MOTA	609	N LEU	82	4.893	-4.048	6.636	1.00 29.86
MOTA	610	CA LEU	82	4.985	-2.738	7.272	1.00 28.10
MOTA	611	CB LEU	82	5.310	-2.871	8.769	1.00 28.19
ATOM	612		82	4.240	-3.477	9.686	1.00 29.26
MOTA	613	CD1 LEU	82	4.674	-3.203	11.133	1.00 28.77
MOTA	614	CD2 LEU	82	2.873	-2.867	9.431	1.00 29.93
MOTA	615	C LEU	82	6.083	-1.916	6.606	1.00 26.82
ATOM	616	O LEU	82	7.241	-2.326	6.582	1.00 26.42
MOTA	617	N PHE	83	5.711	-0.765	6.054	1.00 25.82
	618	•	83	6.673	0.106	5.387	1.00 24.50
ATOM		CB PHE	83	6.008	0.772	4.180	1.00 25.77
MOTA	620		. 83	5.548	-0.209		1.00 27.12
MOTA	621	CD1 PHE			-0.942	3.324	1.00 28.41
ATOM	622	CD2 PHE	83	6.322	-0.447	2.013	1.00 28.63
ATOM	623	CE1 PHE	83		-1.908	2.389	1.00 29.75
ATOM	624	CE2 PHE	83	5.941	-1.408	1.074	1.00 29.85
MOTA	625	CZ PHE	83	4.769	-2.140	1.268	1.00 29.73
MOTA	626	C PHE	83	7.185	1.100	6.382	
ATOM	627	O PHE	83	6.427	1.908	6.910	1.00 22.13
ATOM	628	N LEU	. 84	8.492	1.044	6.628	1.00 21.28
ATOM	629	CA LEU	84	9.144	1.901	7.616	1.00 20.54

FIG.11B-15

ATOM	630	СВ	LEU	84	9.713	1.040	8.745	1 00 10 57
MOTA	631	CG	LEU	84	8.713	0.013	9.290	1.00 19.57 1.00 18.62
ATOM	632	CD1	LEU	84	9.495	-1.049	10.055	1.00 18.62 1.00 19.29
ATOM	633	CD2	LEU	84	7.671	0.657	10.035	1.00 19.29
ATOM	634	C	LEU	. 84	10.331	2.710	7.085	1.00 19.74
ATOM	635	0	LEU	84	10.912	2.396	6.041	1.00 20.28
ATOM	636	N	GLU	85	10.691	3.746	7.834	
ATOM	637	CA	GLU	85	11.828	4.596	7.502	1.00 19.73 1.00 19.25
ATOM	638	CB	GLU	85	11.983	5.690	8.563	1.00 19.25
ATOM	639	CG	GLU	85	13.227	6.565	8.390	1.00 19.19
ATOM	640	CD	GLU	85	13.164	7.676	9.440	1.00 19.09
ATOM	641	0E1	GLU	85	13.955	8.637	9.305	1.00 20.12
ATOM	642	0E2	GLU	85	12.341	7.578		1.00 20.13
ATOM	643	C	GLU	85	13.105	3.768	7.474	1.00 19.30
ATOM	644	. 0	GLU	85	13.454	3.115	8.461	1.00 19.30
ATOM	645	N	TYR	86	13.806	3.775		1.00 18.76
ATOM	646	CA	TYR	8 6	15.037	3.021	6.249	1.00 18.51
ATOM	647	CB	TYR	86	15.406	2.799	4.782	1.00 19.67
ATOM	648	CG	TYR	86	16.774	2.195	4.610	1.00 20.99
ATOM	649	CD1	TYR	86	17.106	0.992	5.233	1.00 21.68
ATOM	650	CE1		8 6 ´	18.372	0.434	5.091	1.00 22.49
ATOM	651		TYR	86	17.747	2.827		1.00 21.14
ATOM	652		TYR	86	19.019	2.272	3.682	1.00 22.21
ATOM	653		TYR	86	19.321	1.082	4.318	1.00 23.27
ATOM	654	OH	TYR	86 .	20.585	0.548	4.216	1.00 25.97
MOTA	655	С	TYR	86 _	16.167	3.769	6.953	1.00 18.53
ATOM	656	0	TYR	86	16.444	4.927	6.631	1.00 17.92
ATOM	657	N	CYS	87	16.797	3.110	7.926	1.00 18.60
ATOM	658	CA	CYS	87	17.904	3.705	8.678	1.00 19.08
ATOM	659	CB	CYS	87	17.697	3.474	10.187	1.00 18.77
MOTA	660	SG	CYS	87	16.171	4.310	10.710	1.00 18.39
ATOM	661		CYS	87	19.193	3.058	8.186	1.00 18.78
ATOM	662	0	CYS	87	19.571	1.968	8.626	1.00 19.02
MOTA	663	N	SER	88	19.879	3.739	7.271	
ATOM	664	CA	SER	88	21.098	3.200	6.687	1.00 20.13
ATOM	665	CB	SER	88 :	21.508	4.021	5.458	1.00 20.76
ATOM	666	OG	SER	88	21.898	5.331	5.835	1.00 21.97
ATOM	667	C	SER	88	22.308	3.098	7.584	1.00 20.67
MOTA	668	0	SER	88	23.273	2.419	7.240	1.00 21.00
ATOM	669	N	GLY	89	22.263	3.758	8.739	1.00 20.40
MOTA	670	CA	GLY	89	23.392	3.718	9.648	1.00 20.92
MOTA	671	С	GLY	89	23.476	2.498	10.544	1.00 20.54

FIG.11B-16

						•			
	ATOM	672	0 -	GLY	89	24.443	2.343	11.285	1.00 21.32
	MOTA	673	N	GLY	90	22.465	1.636	10.497	1.00 20.52
	MOTA	674	CA	GLY	90	22.495	0.435	11.308	1.00 20.74
	MOTA	675	C	GLY	90	22.057	0.669	12.739	1.00 19.72
	ATOM	676	0	GLY	90	21.393		13.041	1.00 19.15
	ATOM	677	N	GLU	91	22.454	-0.243	13.618	1.00 19.25
	MOTA	678	CA	GLU	91	22.095	-0.175	15.032	1.00 18.39
	MOTA	679	CB	GLU	91	21.985	-1.580	15.616	1.00 19.43
	ATOM	680	CG	GLU	91	20.935	-2.428	14.909	1.00 20.97
	MOTA	681	CD	GLU	91	20.884	-3.864	15.432	1.00 21.99
	MOTA	682	√ 0E1	GLU	91	20.081	-4.642	14.863	1.00 23.57
	ATOM	683	0E2	GLU	91	21.624	-4.182	16.387	1.00 20.79
	ATOM	684	C	GLU	91	23.102	0.554	15.861	
	MOTA	685	0	GLU	91	24.289		15.549	1.00 18.50
	ATOM	686	. N	LEU	92	22.628	1.188	16.931	1.00 16.35
	MOTA	687	CA	LEU	92	23.507	1.908		
	ATOM	688	CB	LEU	92	22.684	2.598	18.945	1.00 15.21
	ATOM	689	CG	LEU	92	23.525	3.275	20.041	1.00 13.57
•	ATOM	690		LEU		24.312	4.465	19.512	1.00 14.82
	ATOM	691		LEU		22.545	3.710	21.139	1.00 14.32
:	ATOM	692	C	LEU	92	24.417	0.890	18.448	1.00 17.07
	ATOM	693	0	LEU	92	25.559		18.784	1.00 16.10
•	ATOM	694		PHE	93	23.918	-0.342	18.552	1.00 17.48
	ATOM		CA			24.678	-1.438	19.121	1.00 20.02
٠	ATOM		CB		93	23.888	-2.751	18.999	1.00 21.58
	ATOM		CG	PHE		120	-3.956	19.521	1.00 23.36
	ATOM		CD1			25.553	-4.628	18.721	1.00 23.91
	ATOM	699	-	PHE		24.420	-4.402	20.822	1.00 24.46
	MOTA	700		PHE		26.261	-5.730	19.212	1.00 25.32
	MOTA	701	CE2		93	25.124	-5.506	21.328	1.00 25.27
	ATOM			PHE	93		•	20.522	1.00 24.89
		703			93	26.039		18.425	1.00 21.14
	ATOM		-0	PHE	93		-1.856		1.00 20.61
	ATOM	705	N		94	26.051	-1.450	17.104	1.00 21.54
	ATOM	706	CA	ASP	94		-1.614		1.00 22.83
	ATOM	707	CB	ASP	94	26.908	-1.933	14.857	1.00 24.16
	ATOM	708	CG	ASP	94	26.277	-3.346	14.811	1.00 25.84
	ATOM	•		ASP	94	25.502	-3.688	13.893	1.00 29.46
	ATOM	710		ASP	94	26.543	-4.189	15.686	1.00 26.49
	HOTA	711	C	ASP	94	28.249	-0.425	16.407	1.00 22.49
	MOTA	712	0	ASP.	94	29.365	-0.497	15.896	1.00 23.70
	MOTA	713	N	ARG	95	27.839	0.645	17.084	1.00 21.43

FIG.11B-17

MOTA	714	CA	ARG	95	28.685	1.826	17.250	1.00 20.91
ATOM	715	CB	ARG	95	27.837	3.099	17.129	1.00 23.48
MOTA	716	CG	ARG	95	27.411	3.256	15.674	1.00 26.51
MOTA	717	CD	ARG	95	28.661	3.755	14.919	1.00 28.94
MOTA	718	NE	ARG	95	29.128	5.018	15.492	1.00 32.20
MOTA	719	CZ	ARG	95	28.577	6.203	15.239	1.00 33.03
MOTA	720	NH1	ARG	95	27.544	6.292	14.407	1.00 35.34
MOTA	721	NH2	ARG	95	29.038	7.291	15.836	1.00 33.60
MOTA	722	C	ARG	95	29.378	1.815	18.636	1.00 20.22
MOTA	723	0	ARG	95	30.171	2.706	18.957	1.00 20.16
MOTA	724	N	ILE	96	29.051	0.802	19.435	1.00 19.47
MOTA	725	CA	ILE	96	29.605	0.640	20.771	1.00 18.75
MOTA	726	CB	ILE	96	28.532	0.091	21.721.	1.00 18.34
MOTA	727	CG2	ILE	.96	29.123	-0.162	23.104	1.00 18.88
MOTA	728	CG1	ILE	96	27.371	1.085	21.777	1.00 17.58
MOTA	729	CD1	ILE	96	26.167	0.580	22.596	
MOTA	730	C	ILE	96	30.775	-0.298	20.702	1.00 19.98
MOTA	731	0	ILE	96	30.609	-1.486	20.427	1.00 19.77
MOTA	732	N	GLU	97	31.968	0.230	20.943	1.00 19.99
MOTA	733	CA	GLU	97	33.168	-0.597	20.886	1.00 22.20
MOTA	734	CB	GLU	97	34.383	0.292	20.633	1.00 24.46
MOTA	735	CG	GLU	97	34.631	1.057	19.276	1.00 28.88
MOTA	736	CD	GLU	97	33.720	2.218	18.832	1.00 31.18
MOTA	737	0E1		97	33.585	3.250	19.536	1.00 32.29
MOTA	738		GLU	97	33.142	2.070	17.730	1.00 33.32
MOTA	739	С	GLU	97	33.307	-1.427	22.185	1.00 21.17
MOTA	740	0	GLU	97	33.320	-0.886	23.289	1.00 21.38
MOTA	741	N	PRO	98	33.391	-2.757	22.055	1.00 21.50
MOTA	742	CD	PR0	98	33.282	-3.558	20.817	1.00 21.52
ATOM	743	CA	PR0	98	33.519	-3.622	23.231	1.00 21.75
MOTA	744	CB	PRO	98	33.765	-4.998	22.611	1.00 21.99
MOTA	745	•	PRO	98	32.982	-4.935	21.349	1.00 22.40
MOTA	746	C	PRO	98 .	34.593	-3.186	24.219	1.00 22.45
MOTA	747	0	PR ₀	98	35.722	-2.885	23.827	1.00 22.61
MOTA.	748	N	ASP	99	34.212	-3.168	25.495	1.00 23.48
ATOM	749	CA	ASP	99	35.072	-2.804	26.616	1.00 24.51
MOTA	750	CB	ASP	99	36.323	-3.695	26.634	1.00 27.46
MOTA	751		ASP	99	36.003	-5.182	26.423	1.00 30.42
MOTA	752		ASP	99	35.439	-5.526	25.362	1.00 32.42
ATOM	753	OD2	ASP	99	36.309	-6.023	27.298	1.00 32.91
MOTA	754	C	ASP	99	35.524	-1.341	26.625	1.00 23.52
ATOM	755	0	ASP	99	36.266	-0.917		1.00 23.73

ATOM	756	\1 *1 =					
ATOM	756	N ILE	100	35.082	-0.561	25.650	1.00 22.38
ATOM	757	CA ILE	100	35.490	0.828	25.594	1.00 21.76
ATOM	758	CB ILE	100	36.493	1.045	24.440	1.00 23.97
ATOM	759	CG2 ILE	100	37.824	0.408	24.782	1.00 24.47
ATOM	760	CG1 ILE	100	36.017	0.329	23.181	1.00 25.90
ATOM	761	CD1 ILE	100	37.095	0.266	22.055	1.00 28.42
ATOM	762	C ILE	100	34.351	1.797	25.504	1.00 20.24
MOTA	763	0 ILE	100	34.340	2.797	26.212	1.00 19.97
ATOM	764	N GLY	101	33.389	1.512	24.637	1.00 18.48
MOTA	765	CA GLY	101	32.249	2.405	24.481	1.00 16.99
ATOM	766		101	32.418	3.264	23.241	. *
ATOM	767	O GLY	101	32.595			1.00 17.27
ATOM	768	N MET	102	32.324	4.581		1.00 16.05
ATOM	769	CA MET	102	32.483		22.335	1.00 15.12
ATOM	770	CB MET	102	31.181	5.702		1.00 15.01
ATOM	771	CG MET	102	30.080		22.316	1.00 14.98
ATOM	772	SD MET	102	28.559	6.611		1.00 14.69
ATOM	773	CE MET	102	28.049	4.872		1.00 14.44
ATOM	774	C MET	102	32.834	6.921	22.981	1.00 14.62
ATOM	775	O MET	102	32.713	7.100	24.202	1.00 13.88
ATOM	776	N PRO	103	33.264	7.894	22.171	
ATOM	777	CD PRO	103	33.526	7.844	20.723	1.00 14.81
ATOM	778	CA PRO	103	33.609	9.213	22.715	1.00 14.86
ATOM	779	CB PRO	103	33.984	10.003		1.00 15.81
ATOM	780	CG PRO	103	34.530	8.944	20.559	1.00 16.07
ATOM	781	C PRO	103	32.435		23.479	1.00 15.44
MOTA	782	O PRO	103	31.308	9.789	22.994	1.00 14.56
ATOM	783	N GLU	104		10.351		1.00 14.50
ATOM	784		104		10.948		1.00 15.05
ATOM	785	CB GLU	104	32.263		26.727	1.00 15.48
MOTA	786	CG GLU	104	31.299		27.906	1.00 15.12
ATOM	787	CD GLU	104	32.003			1.00 17.77
ATOM	788	OE1 GLU	104			29.576	1.00 17.77
ATOM	789	OE2 GLU	104	32.848		29.737	
	790	C GLU	104	30.748		24.731	
ATOM	791	O GLU	104	29.533	11.998	24.731	1.00 16.01
ATOM	792	N PRO	105		12.790		1.00 15.90
ATOM	793	CD PRO	105	32.740	13.149	23.840	1.00 16.31
ATOM	794	CA PRO	105			23.642	1.00 17.39
ATOM	795	CB PRO	105	31.427	13.732	23.140	1.00 16.34
ATOM	796	CG PRO	105	_	14.609	22.360	1.00 17.22
ATOM	797	C PRO	105		14.618	23.260	1.00 17.15
ווט ו ר	171	C FRU	TOO	29.418	12.999	22.282	1.00 15.69

FIG.11B-19

MOTA	798	0	PRO	105	28.262	13.414	22.179	1.00 15.94
MOTA	79 9	N	ASP	106	29.846		21.651	1.00 15.00
MOTA	800	CA	ASP	106	28.946		20.810	1.00 15.05
MOTA	801	CB	ASP	106	29.695		20.070	1.00 17.35
ATOM	802	CG	ASP	106	30.678		19.027	1.00 20.16
MOTA	803	OD1	ASP	106	30.627	· · · · · ·	18.686	1.00 23.79
ATOM	804	0D2	ASP	106	31.495		18.541	1.00 24.67
MOTA	805	С	ASP.	106	27.863		21.654	1.00 14.46
ATOM .	806	0	ASP	106	26.696	_	21.240	
MOTA	807	N	ALA -	107	28.249		22.816	1.00 13.14
MOTA	808	CA	ALA	107	27.284		23.692	1.00 12.29
MOTA	809	CB	ALA ·	107	27.990		24.900	1.00 11.81
TATOM	810	C	ALA	107	26.256		24.177	1.00 12.80
ATOM	811	0	ALA	107	25.065	,	24.262	1.00 11.19
MOTA	812	N	GLN	108	26.735	•	24.478	1.00 12.72
ATOM	813	CA	GLN	108	25.838	and the second s	24.964	**
ATOM	814	CB	GLN	108	26.648	•	25.395	1.00 13.39
ATOM	815	CG	GLN	108	25.730		26.208	1.00 13.91
MOTA	816	CD	GLN	108	26.315		26.290	1.00 14.73
MOTA	817	0E1	GLN	108	26.409		27.380	1.00 18.02
MOTA	818	NE2	GLN	108	26.690		25.142	1.00 13.66
MOTA	819	C	GLN.	108	24.828		23.886	1.00 12.49
MOTA	820	0	GLN	108	23.643		24.163	1.00 12.86
ATOM :	821	N	ARG	109	25.298		22.652	
ATOM	822	CA	ARG	109	24.412		21.544	1.00 12.93
MOTA	823	CB	ARG	109	25.242		20.270	
MOTA	824	CG	ARG .	109	24.424	13.899	18.967	1.00 17.36
ATOM	825	CD	ARG	109	25.431	14.120	17.816	1.00 20.06
ATOM	826	NE T	ARG	109	26.088	12.870	17.433	1.00 25.24
MOTA	827	CZ	ARG	109	25.498	11.902	16.732	1.00 25.26
MOTA	828	NH1	ARG	109	24.251	12.039	16.331	1.00 24.49
ATOM	829	NH2	ARG	109	26.157	10.787	16.442	1.00 29.46
MOTA	830	C	ARG	109	23.334		21.342	
MOTA	831	0	ARG	109	22.153			
MOTA	832	N	PHE	110	23.742	11.154	21.345	
MOTA	833	CA	PHE	110	22.778		21.174	
ATOM .	834	CB	PHE	110	23.453			1.00 11.45
MOTA	835	CG	PHE	110	24.187	8.462	19.801	1.00 12.74
MOTA	836	CD1	PHE	110	23.586		18.567	1.00 12.74
ATOM	837	CD2	PHE	110	25.470	_		1.00 12.74
MOTA	838	CE1	PHE	110	24.255			1.00 13.26
MOTA	839	CE2	PHE	110	. 26.142			1.00 14.73
						-		

ATOM	840	CZ PHE	110	25.539	7 000	77 407	1 00 12 07
ATOM	841	C PHE	110	21.819	7.899 10.031	17.401	1.00 13.07
	842	O PHE	110	20.631	:	22.356	1.00 11.04
MOTA					9.735	22.192	1.00 10.10
MOTA	843		111	22.325	10.298	23.558	1.00 10.21
ATOM	844	CA PHE	111	21.455	10.278	24.729	1.00 10.84
ATOM	845	CB PHE	111	22.279	10.392	26.022	1.00 10.93
MOTA	846	CG PHE	111	21.483	10.099	27.265	1.00 11.01
ATOM	847	OD L TITL	111		8.793	27.569	
MOTA	848	CD2 PHE	111	21.087		28.111	1.00 11.89
MOTA		CE1 PHE	111	20.298		28.710	1.00 12.71
MOTA		CE2 PHE		20.307	10.888	29.235	1.00 12.60
MOTA		CZ PHE		19.907		29.536	
MOTA	852		111		11.414	24.651	
MOTA	853	O PHE	111				1.00 10.73
ATOM	854	N HIS	112		12.560		1.00 11.50
MOTA	855	CA HIS	112	19.908	13.678	23.986	1.00 12.31
MOTA	856	CB HIS	· 112 ·	20.562	14.893	23.322	1.00 12.79
MOTA	857	CG HIS	112	21.594	15.584	24.158	1.00 14.79
MOTA	858	CD2 HIS	112	22.655	16.344	23.797	1.00 14.31
MOTA	859	ND1 HIS	112	21.544	15.626	25.534	1.00 15.64
MOTA	860	CE1 HIS	112	22.523	16.389	25.987	1.00 12.99
MOTA	861	NE2 HIS	112	23.212	16.838	24.951	1.00 17.59
MOTA	862	C HIS	112	18.788	13.282	23.019	1.00 12.56
MOTA	863	0 HIS	112	17.608	13.540	23.278	1.00 13.20
MOTA	864	N GLN	113	19.179	12.659	21.906	1.00 12.60
NOTA	865	CA GLN	113	18.226	12.236	20.881	1.00 12.31
MOTA	866	CB GLN	113	18.967	11.756	19.622	1.00 12.91
MOTA	867	CG GLN	113	19.661	12.997	18.985	1.00 13.65
MOTA	868	CD GLN	113	20.372	12.578	17.695	1.00 17.58
MOTA	869	OE1 GLN	113	20.322	11.416	17.302	1.00 20.92
MOTA	870	NE2 GLN	113	21.037	13.530	17.037	1.00 17.64
MOTA				17.338	11.160	21.406	1.00 12.18
MOTA	872						
MOTA	873	N LEU	114	17.906	10.267	22.209	1.00 10.60
MOTA	874	CA LEU				22.804	1.00 10.15
MOTA	875	CB LEU					1.00 9.54
MOTA	876			17.324		24.451	
MOTA	877	CD1 LEU			6.276	23.565	1.00 9.65
ATOM	878	CD2 LEU		18.382	6.296		
ATOM	879	C LEU					•
MOTA	880	O LEU					
ATOM	881		115		10.763		
711011				20.100	10.700		2.00 J.JL

FIG.11B-21

							-		
ATOM	882	CA	MET	115	15	. 485	11.403	25.489	1.00 11.27
MOTA	883	CB	MET	115	16	5.162	12.439	26.386	1.00 12.20
MOTA	884	CG	MET	115	17	7.001	11.884	27.520	1.00 12.71
MOTA	885	SD	MET	115	16	5.069	10.850	28.678	1.00 15.32
MOTA	886	CE	MET	115	16	5.509	9.262	27.927	1.00 10.45
MOTA	887	С	MET	115	14	1.379	12.124	24.719	1.00 11.42
MOTA	888	0	MET	115	13	3.218	12.126		1.00 12.59
ATOM	889	N	ALA	116		1.741	12.739		
MOTA	890	CA	ALA	116		3.762	13.454		1.00 11.80
ATOM	891	CB	ALA	116		4.458	14.124		1.00 12.68
ATOM	892	C	ALA	116		2.697	12.456		1.00 12.52
ATOM	893	Ō	ALA	116		1.496	12.737		1.00 12.58
ATOM	894	N .	GLY	117			. 11.299		1.00 10.70
MOTA	895	CA	GLY	117		2.236	10.276		
ATOM	896	C	GLY	117		1.375	9.700		1.00 11.35
MOTA	897	0	GLY	117		0.176	9.490		
ATOM	898	N	VAL	118		1.976	9.441	_	1.00 11.46
ATOM	899	CA	VAL	118		1.221	8.877		1.00 10.70
ATOM	900	CB	VAL	118		2.191	8.367		1.00 10.80
ATOM	901		VAL	118		1.423	7.893		1.00 11.15
ATOM	902		VAL	118.		3.005	7.230		1.00 10.87
MOTA	903	C	VAL	118		0.199	9.886		1.00 10.07
ATOM	904	Ŏ	VAL	118		9.043	9.514		1.00 12.27
ATOM	905	N	VAL	119		0.619			· · · · · · · · · · · · · · · · · · ·
MOTA	906	CA	VAL	119		9.718	12.200		1.00 13.60
MOTA	907	CB	VAL	119		0.376	13.614		•
ATOM	908	CG1		119		9.310	14.696		1.00 14.89
ATOM	909		VAL	119		1.385	13.763		1.00 14.60
ATOM	910	C	VAL	119		8.506	12.256		1.00 13.79
ATOM	911	0	VAL	119		7.355	12.380		1.00 14.60
MOTA	912	N	TYR	120		8.773	12.159		
ATOM	913·		TYR	120		7.687			1.00 13.72
ATOM	914	CB	TYR	120		8.243	12.109		
ATOM	915		TYR	120			11.900	· · · · · · · · · · · · · · · · · · ·	1.00 15.90
ATOM	916		TYR	120		6.309			
ATOM	917		TYR	120		5.250			· ·
ATOM	918		TYR	120		6.924			
	919		TYR	120					
ATOM	•					5.869			•
ATOM	920	CZ	TYR	120		5.038		•	1.00 18.22
ATOM	921	OH	TYR	120		3.998			
ATOM	922	C	TYR	120		6.705			1.00 13.65
ATOM	923	0	TYR	120		5.481	11.193	23.015	1.00 13.76

ATOM	924	N LEU	121	7.245	9.786	22 060	1 00 11 07
ATOM	925	CA LEU	121	6.407	8.610	22.968 23.155	1.00 11.97 1.00 11.14
ATOM	926	CB LEU	121	7.262	7.337	23.236	1.00 11.14
ATOM	927	CG LEU	121	8.001	6.961	21.937	1.00 10.46
ATOM	928	CD1 LEU	121	8.830	5.695	22.199	1.00 9.60
ATOM	929	CD2 LEU	121	7.039	6.749	20.774	1.00 12.56
ATOM	930	C LEU	121	5.576		24.452	1.00 11.38
ATOM	931	O LEU	121		8.479		1.00 10.61
ATOM	932	N HIS	122	6.234		25.553	1.00 10.61
ATOM	933	CA HIS	122	5.524	9.194	26.820	1.00 11.84
ATOM	934	CB HIS	122	6.528	9.447		1.00 12.54
ATOM	935	CG HIS	:	7.381	8.255	28.262	
ATOM	936	CD2 HIS	•	7.382		27.747	
MOTA		ND1 HIS	122	8.348		29.248	
ATOM		CE1 HIS	122	8.905	7.070	29.328	1.00 11.31
ATOM		NE2 HIS	and the second s	8.335		28.431	1.00 10.75
ATOM		C HIS			10.255		1.00 12.85
ATOM	941	O HIS	122	3.391	10.127	ed and a second second	1.00 13.14
MOTA	942	N GLY	123	4.724	11.291	1 15 15 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	1.00 13.41
MOTA	943	CA GLY	123	3.767		25.838	
MOTA	944	C GLY	123	2.469	11.927	25.198	1.00 16.34
MOTA	945	O GLY	123	1.398	12.472		1.00 17.30
MOTA	946	N ILE	124	2.555	10.946	24.305	1.00 15.33
MOTA	947	CA ILE	124	1.373	10.429	23.625	1.00 16.62
MOTA	948	CB ILE	124	1.660	10.125	22.128	1.00 19.17
MOTA	949	CG2 ILE	124	2.685	9.029	21.991	1.00 20.31
MOTA	950	CG1 ILE	124	0.365		21.423	1.00 21.41
MOTA	951	CD1 ILE	124		10.819	21.404	1.00 24.63
MOTA	952		124	0.840	9.193	24.325	1.00 15.39
MOTA		O ILE	124	-0.067		23.821	1.00 16.38
MOTA		N GLY	125	1.418	8.873	25.481	1.00 14.25
MOTA	955			•		26.270	
ATOM	956	C GLY	125				1.00 14.37
MOTA	957	O GLY	125	0.787	5.364		
MOTA	958	N ILE	126		5.304		
MOTA	959	CA ILE	126			24.644	1.00 14.92
MOTA	960	CB ILE	126	3.259		23.088	1.00 17.26
MOTA	961	CG2 ILE			5.480	22.346	1.00 18.55
MOTA	962	CG1 ILE	126	4.559	5.667		1.00 21.12
MOTA	963	CD1 ILE	126	5.816	4.689		1.00 23.73
ATOM	964	C ILE	126	4.355	•	25.332	
MOTA	965	O ILE	. 126	5.209	5.554	25.532	1.00 12.38

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MOTA	966	N	THR	127	4.514	3.416	25.725	1.00 12.02
MOTA	967	CA	THR	127	5.787	2.991	26.304	1.00 12.59
MOTA	968	CB	THR	127	5.630	2.446	27.729	1.00 13.49
MOTA	9 69	0G1	THR	127	5.288	3.516	28.613	1.00 21.14
MOTA	970	CG2	THR	127	6.923	1.830	28.190	1.00 8.53
MOTA	971	C	THR	127	6.362	1.951	25.381	1.00 11.65
MOTA	972	0	THR	127	5.646	1.065	24.907	1.00 12.97
MOTA	973	N	HIS	128	7.665	2.054	25.103	1.00 12.24
MOTA	974	CA	HIS	128	8.321	1.123	24.187	1.00 12.15
MOTA	975	CB	HIS	128	9.648	1.736	23.711	1.00 11.49
MOTA	976	CG	HIS	128	10.375	0.904	22.699	1.00 11.68
MOTA	977	CD2	HIS	128	10.471	1.012		1.00 11.46
MOTA	978	ND1	HIS	128	11.119		23.050	1.00 11.90
MOTA	979	CE1	HIS	128	11.641	-0.741	21.961	1.00 12.68
MOTA	980	NE2	HIS	128	11.262	-0.025	20.915	1.00 12.57
MOTA	981	C	HIS	128	8.517	-0.242	24.817	1.00 12.15
ATOM	982	0	HIS	128	8.260	-1.275	24.192	1.00 11.72
MOTA	983	N	ARG	129	8.968	-0.236	26.070	1.00 11.07
ATOM	984	CA	ARG	129	9.191	-1.462	26.849	1.00 11.47
MOTA	985	CB	ARG	129	7.931	-2.343	26.858	1.00 12.32
MOTA	986	CG	ARG	129	6.807	-1.547	27.487	1.00 12.84
MOTA	987	CD	ARG	129	5.709	-2.441	28.097	1.00 12.85
ATOM	988	NE	ARG	129	4.988	-3.179	27.067	1.00 12.16
ATOM	989	CZ	ARG	129	3.911	-3.911	27.316	
MOTA	990		ARG	129	3.446	-3.991	28.565	1.00 13.82
MOTA	991	NH2	ARG	129	3.304	-4.553	26.317	1.00 14.27
MOTA	992	C	ARG	129	10.380	-2.359	26.501	1.00 12.13
MOTA	993	0	ARG	129	10.592	-3.375	27.159	1.00 11.76
MOTA	994	•	ASP	130	11.162	-1.999	25.488	1.00 10.41
MOTA	995	CA	ASP	130	12.332	-2.823	25.147	1.00 11.17
MOTA	996	CB	ASP	130	11.914	-3.937	24.157	1.00 11.51
MOTA		CG	ASP ·	130	12.950	-5.082	24.156	1.00 13.34
MOTA	998		ASP.	130	12.988	-5.895	23.184	1.00 13.30
ATOM	999		ASP	130 🐇	13.732	-5.178	25.127	1.00 13.23
MOTA	1000	C.	ASP	130	13.442	-1.969	24.584	1.00 10.24
MOTA	1001	0	ASP	130	14.048	-2.300	23.564	1.00 10.99
MOTA	1002	N "	ILE	131	13.735	-0.848	25.245	1.00 9.92
ATOM	1003	CA	ILE	131	14.787	0.047	24.763	1.00 10.16
MOTA	1004	CB	ILE	131	14.705	1.408	25.463	1.00 10.21
MOTA	1005	CG2		131	15.892		25.040	1.00 10.96
ATOM	1006	CG1		131	13.350	2.041	25.136.	1.00 10.91
MOTA	1007	CD1	ILE	131	13.075	3.389	25.902	1.00 12.36

FIG.11B-24

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MOTA	1008	С	ILE	131	16.152	-0.580	25.017	1.00 10.07
MOTA	1009	0	ILE	131	16.449	-0.979	26.134	1.00 11.34
MOTA	1010	N	LYS	132	16.969	-0.643	23.970	1.00 10.08
MOTA	1011	CA	LYS	132	18.314	-1.214	24.029	1.00 10.81
MOTA	1012	CB	LYS	132	18.256	-2.741	24.204	1.00 10.84
MOTA	1013		LYS	132	17.367	-3.496	23.156	1.00 11.39
MOTA	1014	CD	LYS	132	17.443	-5.018	23.486	1.00 13.71
MOTA	1015	CE	LYS	132	16.477	-5.789	22.554	1.00 13.34
MOTA	1016	NZ	LYS	132	16.456	-7.271	22.822	1.00 13.78
MOTA	1017	C	LYS	132	18.974	-0.793	22.721	1.00 11.06
MOTA	1018	0	LYS	132	18.285	-0.412	21.767	1.00 12.07
MOTA	1019	N	PRO	133	20.307	-0.857	22.651	1.00 12.09
MOTA	1020	CD	PR0	133	21.235	-1.277	23.717	1.00 11.84
MOTA	1021	CA	PR0	133	21.040	-0.455	21.442	1.00 12.30
MOTA	1022	CB	PRO	133	22.499	-0.707	21.838	1.00 11.29
MOTA	1023	CG	PRO	133	22.486	-0.504	23.347	1.00 11.82
ATOM	1024	C	PR0	133	20.595	-1.090	20.156	1.00 13.29
MOTA	1025	0	PRO	133	20.667	-0.452		1.00 12.31
ATOM	1026	N	GLU	134	20.120	-2.335		1.00 13.34
ATOM	1027	CA	GLU	134	19.657		19.047	1.00 14.29
ATOM	1028	CB	GLU	134	19.393	-4.512	19.354	1.00 15.51
ATOM	1029	CG	GLU	134	20.601	-5.281		1.00 16.70
ATOM	1030	CD	GLU	134		-5.274		
ATOM	1031		GLU	134	20.556	-4.248		1.00 16.56
ATOM	1032		GLU	134	21.194		21.905	1.00 19.18
ATOM	1033		GLU	134	18.372		18.486	•
ATOM			GLU	134	18.064	•	17.293	
ATOM	1035		ASN	135	17.625	-1.715		1.00 13.51
MOTA	1036		ASN	135	16.367	-1.086		1.00 12.98
MOTA	1037		ASN	135	15.252	-1.396		
MOTA	1038		ASN	135	14.698	-2.831		1.00 13.89
MOTA	1039	*.	ASN					1.00 15.19
MOTA	1040		ASN	135				1.00 13.17
ATOM	1041		ASN	135	16.471		18.760	•
ATOM	1042		ASN	135	15.462			1.00 13.83
ATOM	1043		LEU	136	17.699			
MOTA	1044		LEU	136	•	2.330		1.00 12.25
ATOM	1045		LEU		18.689		19.593	•
MOTA	1046		LEU	136	17.899		20.912	
MOTA	1047		LEU		18.839		-	
ATOM	1048		LEU		16.599		•	
ATOM	1049	C	LEU	136	18779	2.378	17.091	1.00 13.15

FIG.11B-25

MOTA	1050	0	LEU	136		19.957	2.043	17.084	1.00:13.18
MOTA	1051	N	LEU	137		18.124	2.735	15.991	1.00 13.60
MOTA	1052	CA	LEU	137		18.783	2.775	14.692	1.00 14.28
MOTA	1053	CB .	LEU	137		17.814	2.291	13.611	1.00 14.14
MOTA	1054	CG	LEU	137		17.210	0.941	14.025	1.00 15.38
ATOM	1055	CD1	LEU	137		16.280	0.448		1.00 14.24
MOTA	1056	CD2	LEU	137		18.300	-0.074	14.319	1.00 15.67
ATOM	1057	C .	LEU	137		19.287		14.359	1.00 15.19
ATOM	1058	0 .	LEU	137		18.884	5.118	14.952	1.00 15.24
ATOM	1059	N	LEU	138		20.174	4.178	13.372	1.00 16.26
ATOM	1060	CA	LEU	138	٠.	20.779	5.433	12.952	1.00 16.85
MOTA	1061	CB	LEU	138		22.296	.5.385	13.183	1.00 17.95
MOTA	1062	CG	LEU	138	÷, :	22.811	5.144	14.617	1.00 19.40
MOTA	1063	CD1	LEU	138		22.251	6.225	15.531	1.00 19.08
ATOM	1064	CD2	LEU	138		22.399	3.760	15.102	1.00 22.69
MOTA	1065	C	LEU	138		20.534-	5.671	11.461	1.00 17.74
ATOM	1066	0	LEU	138 :		20.604	4.731	10.676	1.00 17.49
ATOM	1067	N	ASP	139		20.236	6.913	11.083	1.00 18.21
MOTA	1068	CA	ASP	139		20.013	7.235	9.673	1.00 20.36
MOTA	1069	CB	ASP	139		18.989	8.371	9.527	1.00 20.49
MOTA	1070	CG	ASP	139		19.372	9.764	9.970	1.00 20.35
MOTA	1071		ASP	139		18.491	10.652	9.888	1.00 23.24
MOTA	1072		ASP	139		20.517	10.002	10.389	1.00 21.14
	1073		ASP.	139		21.345		9.073	1.00 21.49
MOTA	1074	0	ASP	139		22.381		9.709	1.00 20.83
MOTA	1075	N	GLU	140		21.330	8.115		1.00 23.55
MOTA	1076	CA.	GLU	140		22.564	8.511	7.169	1.00 25.49
MOTA	1077	CB	GLU	140		22.282	8.952	5.726	1.00 27.10
MOTA	1078		GLU	140		21.287	10.082	5.469	1.00 30.90
MOTA	1079	CD	GLU	140	-	19.954	9.585		1.00 32.27
MOTA	1080	OE1		140		19.575	8.466	5.572	1.00 34.20
ATOM				•		19.282			1.00 35.13
	1082		GLU	140		23.386		7.867	1.00 25.97
· .	1083		GLU	140					1.00 26.62
MOTA	1084	. N .	ARG	141		22.736	10.444	8.692	1.00 26.36
•	1085		ARG	141				9.408	
MOTA	1086	CB	ARG	141			12.821	9.362	1.00 28:11
MOTA	1087	CG	ARG	141		22.492			1.00 30.79
MOTA	1088		ARG	141			14.853	7.950	1.00 32.84
MOTA	1089		ARG	141			15.861	8.833	1.00 35.77
MOTA	1090	CZ	ARG	141			16.036		1.00 37.04
ATOM	1091	NH1	ARG	141		21.039	15.271	10.681	1.00 36.95

MOTA	1092	NH2 ARG	141	22.564	16.981	10.828	1.00 37.88
ATOM	1093	C ARG	141	23.645	11.156	10.879	1.00 25.65
MOTA	1094	O ARG	141	23.877	12.026	11.720	1.00 25.46
MOTA	1095	N ASP	142	23.579	9.868	11.184	1.00 23.88
MOTA	1096	CA ASP	142	23.755	9.403	12.559	1.00 23.96
ATOM	1097	CB ASP	142	25.128	9.800	13.107	1.00 26.22
ATOM	1098	CG ASP	142	26.194	8.903	12.523	1.00 28.88
ATOM	1099	OD1 ASP	142	25.919	7.694	12.418	1.00 29.81
MOTA	1100	OD2 ASP	142	27.300	9.375	12.182	1.00 31.98
ATOM	1101	C ASP	142	22.677	9.917	13.533	1.00 22.68
MOTA	1102	O ASP	142	22.940	10.092	14.729	1.00 21.46
ATOM	_1103	N ASN	143	21.475	10.172	13.025	1.00 21.05
ATOM	1104	CA ASN	143	20.387	10.603	13.909	1.00 19.86
MOTA	1105	CB ASN	143	19.326	11.401	13.156	1.00 19.72
ATOM	1106	CG ASN	143	19.848	12.766	12.784	1.00 20.89
MOTA	1107	OD1 ASN	143	19.752	13.190	11.621	1.00 23.00
MOTA	1108	ND2 ASN	143	20.404	13.472	13.762	1.00 18.41
ATOM	1109	C ASN	143	19.749			1.00 18.33
ATOM	1110	O ASN	143	19.447	8.437	13.698	1.00 18.35
ATOM	1111	N LEU	144	19.536			
ATOM	1112	CA LEU					1.00 16.09
ATON	1113	CB LEU	144		8.075		1.00 15.65
ATOM		CG LEU				18.855	
ATOM	–	CD1 LEU	144			19.992	
MOTA		CD2 LEU	144		7.585		1.00 15.73
ATON		C LEU	144			16.194	
ATOM	1118	: .	144			16.265	
MOTA	1119				3.7		
ATOM	1120		145			15.744	
MOTA		CB LYS	145			14.277	
ATOM	1122		•	15.953		13.201	1.00 18.61
MOTA	1123		145	· ·			1.00 19.20
ATOM	1124		145	15.797		12.033	
ATOM	1125		145			11.574	•
MOTA	1126			15.126			·
	1127		145			16.329	· ·
MOTA	1128		146	14.166		17.436	1.00 13.48
MOTA	1129	•		13.654			
ATOM	1130		146				
ATOM	1131	•	146			20.070	•
MOTA	1132		146	13.538	_	20.324	•
ATOM	1133	CD1 ILE	146	12.694	6.479	21.403	1.00 13.33

FIG.11B-27

MOTA	1134	C	ILE	146	12.901	3.476	17.301	1.00 13.05
MOTA	1135	0	ILE	146	12.012	3.904	16.559	1.00 12.88
MOTA	1136	N	SER	147	13.238	2.192	17.382	1.00 12.43
MOTA	1137	CA	SER	147	12.681	1.173	16.496	1.00 13.35
MOTA	1138	CB	SER	147	13.822	0.665	15.593	1.00 14.71
ATOM	1139	0G	SER	147	13.489	-0.508	14.855	1.00 15.66
ATOM	1140	C .	SER	147	12.038	-0.011	17.174	1.00 14.00
MOTA	1141	0	SER	147	12.369	-0.344		1.00 12.67
MOTA	1142	N.	ASP	148	11.110	-0.640	16.451	1.00 14.20
MOTA	1143	CA	ASP	148	10.417	-1.855	16.883	1.00 14.80
MOTA	1144	CB	ASP	148	11.453	-2.915	17.282	1.00 16.84
MOTA	1145	CG	ASP	148	10.867	-4.294	17.493	1.00 19.50
MOTA	1146	OD1	ASP	148	11.660	-5.228	17.723	1.00 23.01
MOTA	1147	0D2	ASP	148	9.636	-4.457	17.430	1.00 19.78
ATOM	1148	C	ASP	148	9.426	-1.695	17.980	1.00 15.21
MOTA	1149	0	ASP	148	9.767	-1.794	19.152	1.00 15.83
MOTA	1150	N	PHE	149	8.166	-1.494	17.610	1.00 14.13
MOTA	1151	CA	PHE	149	7.101	-1.309	18.585	1.00 14.87
MOTA	1152	CB	PHE	149	6.252	-0.114	18.143	1.00 15.09
MOTA	1153	CG	PHE	149	6.974	1.187	18.274	1.00 15.25
ATOM	1154	CD1	PHE	149	7.860	1.608	17.292	1.00 15.38
MOTA	1155	CD2	PHE	149	6.844	1.952	19.440	1.00 14.86
MOTA	1156	CE1	PHE	149	8.623	2.776	17.464	1.00 14.94
MOTA	1157	CE2	PHE	149	7.599	3.114	19.621	1.00 14.99
MOTA	1158	CZ	PHE	149	8.487	3.524	18.636	1.00 14.89
MOTA	1159	C	PHE	149	6.304	-2.544	18.811	1.00 15.24
MOTA	1160	0	PHE	149	5.145	-2.484	19.220	1.00 16.36
MOTA	1161	N	GLY	150	6.936	-3.691	18.572	1.00 16.17
ATOM	1162	CA	GLY	150	6.269	-4.970	18.746	1.00 16.47
ATOM	1163	C	GLY	150	5.947	-5.286		1.00 16.83
MOTA	1164	0	GLY	150	5.093	-6.126	20.481	1.00 17.90
MOTA	1165	N	LEU	151	6.621	-4.626	21.127	1.00 15.78
ATOM.	1166	CA		151	6.326	-4.871	22.541	1.00 16.37
MOTA	1167	CB	LEU	151	7.584	-5.298	23.292	1.00 17.64
ATOM	1168		LEU	151	8.078	-6.619	22.700	1.00 19.89
MOTA	1169			151	9.341	-6.989	23.457	1.00 19.60
MOTA	1170	CD2	LEU	151	7.040	-7.730	22.782	1.00 20.88
ATOM	1171	C	LEU	151	5.729	-3.663	23.222	1.00 15.93
ATOM	1172	0	LEU	151	5.392	-3.723	24.405	1.00 16.32
MOTA	1173	N	ALA	152	5.567	-2.577	22.466	1.00 14.33
MOTA	1174	CA	ALA	152	5.020	-1.329	22.994	1.00 14.78
ATOM	1175	CB	ALA	152	5.196	-0.205	21.964	1.00 14.02

FIG.11B-28

ATOM	1176	C .	ala 🗀	152	:	3.559	-1.412	23.400	1.00 14.74
MOTA	1177	0	ALA	152		2.820	-2.287	22.946	1.00 16.44
ATOM	1178	N	THR	153		3.134	-0.498	24.262	1.00 14.29
MOTA	1179	CA	THR	153		1.736	-0.486	24.678	1.00 14.48
MOTA	1180	CB	THR	153	•	1.465	-1.570	25.767	1.00 15.01
MOTA	1181	0G1	THR	153	· · · .	0.052	-1.803	25.857	1.00 15.18
MOTA	1182	CG2	THR	153		1.985	-1.140	27.135	1.00 16.34
ATOM			THR	153	ř.	1.305	0.866	25.134	1.00 14.74
ATOM	1184	0	THR	153		2.124	1.764	25.321	1.00 14.16
ATOM	1185	N	VAL	154		-0.002	1.036	25.283	1.00 15.49
ATOM	1186	CA	VAL	154		-0.574	2.294	25.744	1.00 16.27
ATOM	1187	CB		154	- 1	-2.024	2.446	The second second second	1.00 17.58
MOTA		CG1		154.		-2.721	3.669	25.796	1.00 18.88
ATOM	1189		VAL	154		-1.978	2.573		1.00 18.48
MOTA	1190		VAL	154		-0.584	2.280	27.288	
ATOM	1191	0	VAL	154		-1.096	1.337	27.896	1.00 17.03
ATOM	1192	N		155		0.002		27.917	
ATOM	1193		PHE	155		-0.011	3.348	29.372	1.00 14.65
MOTA	1194	. 1. 77	PHE	155		1.411	3.401		1.00 13.64
MOTA	1195		PHE	155		2.088	4.733	29.830	1.00 12.64
MOTA	1196		PHE	155	1	2.810	5.044	28.675	1.00 11.40
ATOM	1197	CD2	PHE	155		1.984	5.696	30.836	1.00 12.17
ATOM	1198	CE1	PHE	155		3.412		28.520	1.00 12.13
ATOM	1199	CE2	PHE	155		2.592	6.960	30.691	1.00 13.02
MOTA	1200	CZ	PHE	155		3.311	7.254	29.522	1.00 12.14
MOTA	1201	C	PHE	155	ar en en	-0.830	4.566	29.831	1.00 14.29
ATOM	1202	0	PHE	155		-1.041	4.748	31.027	1.00 14.95
ATOM	1203	N	ARG	156		-1.257	5.406	28.888	1.00 14.44
ATOM	1204	CA	ARG	156		-2.082	6.570	29.246	1.00 14.70
ATOM	1205	CB	ARG	156	1. 7	-1.241	7.846	29.410	1.00 15.59
ATOM	1206	CG	ARG	156	· · · · · .	-2.174	8.962	30.089	1.00 17.04
ATOM	1207	CD	A RG	156	.* .	-1.525	10.389	29.970	1.00 18.38
MOTA	1208	NE	ARG	156	·	-0.159	10.425	30.482	1.00 18.17
MOTA	1209	CZ	ARG	156		0.922	10.609	29.719	1.00 18.70
ATOM	1210	NH1	ARG	156		0.795	10.779	28.411	1.00 18.30
MOTA	1211	NH2	ARG	156		2.131	. 10.605	30.265	1.00 19.27
MOTA	1212	C	ARG	156		-3.100	6.807	28.154	1.00 14.58
MOTA	1213	0	ARG	156		-2.753	6.945	26.984	1.00 14.45
MOTA	1214	N	TYR	157		-4.372	6.858	28.541	1.00 14.25
MOTA	1215	CA	TYR	157		-5.441	7.060	27.574	1.00 14.50
MOTA	1216		TYR	157	٠	-6.040	5.715	27.173	1.00 15.15
MOTA	. 1217	CG	TYR ·	157		-6.845	5.770	25.902	1.00 15.19

FIG.11B-29

MOTA	1218	CD1	TYR -	157	-6.219	5.814	24.653	1.00 16.14
MOTA	1219	CE1	TYR	157	-6.965	5.864	23.472	
MOTA	1220	CD2	TYR	157	-8.232	5.780	25.945	1.00 15.85
MOTA	1221	CE2	TYR	157	-8.987	5.832	24.783	1.00 17.07
MOTA	1222	CZ	TYR	157 ·	-8.356	5.875	23.548	1.00 17.48
MOTA	1223	OH	TYR	157	-9.129	5.956	22.403	1.00 17.82
MOTA	1224	C	TYR.	157	-6.507	7.890	28.231	1.00 15.01
ATOM	1225	0	TYR-	157	-6.867	7.632	29.379	1.00 15.42
ATOM	1226	N .	ASN	158	-7.013	8.887	27.505	1.00 16.30
ATOM	1227	CA	ASN	158	-8.033	9.786	28.047	1.00 16.67
MOTA	1228	CB	ASN	158	-9.345	9.023	28.285	1.00 16.49
ATOM	1229	CG	ASN	158	-10.097	8.800	26.961	1.00 15.70
MOTA	1230	OD1 ·	ASN;	. 15 8	-10.988	7.954	26.882	1.00 16.14
ATOM	1231	ND2	ASN	158	-9.741	9.569	25.927	1.00 13.85
ATOM	1232	C	ASN	158	-7.543	10.420	29.336	1.00 18.61
MOTA	1233	0	ASN	158	-8.312	10.640	30.278	1.00 17.84
ATOM	1234	N	ASN	159	-6.242	10.706	29.348	1.00 19.74
MOTA	1235	CA	ASN	159	-5.530	11.321	30.462	1.00 22.33
MOTA	1236	CB	ASN	159	-6.099	12.713	30.758	1.00 24.30
MOTA	1237	CG	ASN	159	-4.976	13.515	31.438	1.00 26.68
MOTA	1238	OD1	ASN	159	-3.879	13.667	30.885	1.00 28.50
MOTA	1239	ND2	ASN	159	-5.249	14.021	32.633	1.00 29.10
MOTA	1240	C .	ASN	159	-5.522	10.478	31.742	1.00 22.20
MOTA	1241	:. 0	ASN	159	-5.259	10.992	32.824	1.00 24.01
ATOM	1242	N	ARG	160	-5.808	9.185	31.621	1.00 21.47
MOTA	1243	CA	ARG	160	-5.803	8.298	32.781	1.00 20.69
MOTA	1244	CB	ARG	160	-7.141	7.559	32.907	1.00 23.48
MOTA	1245	CG	ARG	160	-8.091	8.097	34.011	1.00 28.38
ATOM	1246	CD	ARG	160	-7.621	7.848	35.471	1.00 30.43
MOTA	1247	NE	ARG	160	-8.739	7.883	36.411	1.00 34.64
ATOM	1248	CZ	ARG	160	-9.202	8.978	37.008	1.00 35.54
ATOM	1249		ARG					1.00 36.58
ATOM	1250	NH2	ARG	160	-10.246	8.890	37.829	1.00 37.21
MOTA	1251	C	ARG .	160	-4.668	7.290	32.612	1.00 19.16
ATOM	1252	0	ARG	160	-4.591	6.601	31.602	1.00 18.07
ATOM	1253	N	GLU	161	-3.778	7.225	33.597	1.00 17.48
ATOM	1254	CA	GLU	161	-2.654	6.297	33.530	1.00 17.30
MOTA	1255	CB	GLU	161	-1.528	6.741	34.468	1.00 16.64
MOTA	1256	CG	GLU	161	-0.264		34.412	1.00 16.91
ATOM	1257	CD	GLU	161	0.821	6.223	35.416	1.00 18.42
MOTA	1258	0E1	GLU	161	1.882	5.569	35.377	1.00 16.94
MOTA	1259	0E2	GLU	161	0.606	7.154	36.224	1.00 19.58

MOTA	1260	C	GLU	161	-3.060	4.909	33.903	1.00 17.47
MOTA	1261	0.	GLU:	161	-3.846	4.696	34.836	1.00 17.69
MOTA	1262	N	ARG	162	-2.522	3.941	33.177	1.00 18.18
MOTA	1263	CA	ARG	162	-2.785	2.536	33.425	1.00 18.75
MOTA	1264	CB	ARG	162	-3.133	1.824	32.121	1.00 22.41
MOTA	1265	CG	ARG	162	-3.510	0.361	32.099	1.00 26.57
MOTA	1266	CD	ARG	162	-4.025	0.042	30.639	1.00 29.14
MOTA	1267	NE	ARG	162	-5.085	0.956	30.197	1.00 32.40
MOTA	1268	CZ	ARG	162	-5.832	0.771	29.106	1.00 32.85
ATOM -	1269	NH1	ARG	162	-6.771	1.651	28.776	1.00 33.54
MOTA	1270	NH2	ARG	162	-5.649	-0.301	28.346	1.00 33.73
ATOM	1271	·C	ARG	162	-1.485	1.899	33.950	1.00 17.95
MOTA	1272	0 .	ARG .	162	-0.453	2.015		and the second s
ATOM	1273	N .	LEU	163	-1.532		the second of the second	
ATOM	1274	CA	LEU	163		0.593		1.00 15.46
ATOM	1275	CB	LEU	163			37.117	1.00 16.31
ATOM	1276	CG		N	-0.758			1.00 16.62
MOTA	1277	January Toronto	LEU	163	-1.147	1.014		1.00 17.07
ATOM	1278	1.00	LEU	163	0.467	2.387		
MOTA	1279	C		163		-0.685		Part of the second seco
ATON	1280	0	LEU	163	-1.077	-1.349		1.00 16.39
ATON	1281	N	LEU	164	1.147	-1.031		
ATOM	1282	CA	LEU	164	1.465	-2.231		
ATOM	1283		LEU		are a second of the second of	-1.950	and the second of the second	1.00 15.48
ATOM	1284	CG		164		-0.814		
MOTA	1285		LEU	164	3.528		•	
MOTA		CD2		164		* * * * * * * * * * * * * * * * * * * *		
ATOM	1287		LEU	164	1.811	-3.403		1.00 14.77
MOTA	1288		LEU	164			•.	
MOTA	1289		ASN	165			34.197	1.00 15.82
MOTA	1290		ASN	165	2.017		34.962	
	1291			-				1.00 18.54
MOTA	1292			165				1.00 18.94
ATOM	1293			165		-6.122	•	
MOTA	1294			165		-7.887		
ATOM	1295		ASN	165				
MOTA	1296		ASN	165		-7.887		
MOTA	1297		LYS	166	2.701			1.00 18.68
MOTA	1298		LYS	166	3.297		31.862	•
ATOM	1299		LYS	166	2.916	-7.359	•	
MOTA	1300		LYS	166	3.530			
MOTA	1301	CD	LYS	166	3.333	-7.673	27.802	1.00 26.84

FIG.11B-31

ATOM	1302	CE	LYS	166	,	3.949	-8.340	26.520	1.00 27.03
ATOM	1303	NZ	LYS	166		5.449	-8.227	26.415	1.00 28.54
ATOM -	1304	C	LYS	166		4.794	-7.686	31.950	1.00 20.70
MOTA	1305	0	LYS	166		5.447	-6.639	31.963	1.00 18.98
MOTA	1306	N	MET	167		5.355	-8.886	32.019	1.00 21.49
MOTA	1307	CA	MET	167		6.800	-9.013	32.071	1.00 23.10
MOTA	1308	CB	MET	167		7.203	-10.281	32.863	1.00 26.20
MOTA	1309	CG	MET	167	٠.	7.427	-10.090	34.463	1.00 29.77
ATOM	1310	SD	MET	167	٠.	7.743	-11.672	35.352	1.00 36.62
MOTA	1311	CE	MET	167		6.109	-12.412		1.00 34.01
MOTA	1312	C	MET	167		7.298	-9.031	30.615	1.00 22.75
ATOM	1313	0	MET	167		6.861	-9.837	29.789	1.00 22.16
MOTA	1314	.∴N:	CYS	· 168		8.169	-8.083	30.292	1.00 21.44
ATOM	1315	CA	CYS	168	••	8.750	-8.011	28.963	1.00 20.77
MOTA	1316	CB	CYS	. 168		7.754	-7.523	27.926	1.00 22.35
MOTA	1317	SG	CYS	168	• :	6.960	-5.964	28.305	1.00 26.07
MOTA	1318	C	CYS	168		9.915	-7.126	28.970	1.00 18.90
MOTA	1319	0	CYS	168		10.132	-6.357	29.914	1.00 17.73
MOTA	1320	N	GLY	169		10.696	-7.219	27.903	1.00 17.73
MOTA	1321	CA	GLY	169		11.908	-6.437	27.812	1.00 15.14
MOTA	1322	C	GLY	169		13.074	-7.384	27.579	1.00 15.05
MOTA	1323	0	GLY	169		12.889	-8.485	27.043	1.00 14.39
MOTA	1324	N	THR	170		14.264	-6.957	27.990	1.00 12.82
MOTA	1325	CA	THR	170		15.498	-7.726	27.817	1.00 13.82
MOTA	1326	CB	THR	170		16.278	-7.119	26.624	1.00 12.90
MOTA	1327		THR	170	•	15.476	-7.208	25.432	1.00 13.36
MOTA	1328		THR	170		17.582	-7.853	26.399	1.00 14.59
ATOM	1329	C,	THR	170		16.183	-7.607	29.174	1.00 13.27
MOTA	1330	0	THR	170	• •	16.504	-6.502	29.615	1.00 13.03
MOTA	1331	N	LEU	171	·	16.412	-8.744	29.830	1.00 13.36
MOTA	1332	CA	LEU	171		16.961	-8.761	31.187	1.00 14.06
ATOM	1333	CB	LEU	171	: •	17.427	-10.190	31.522	1.00 15.16
MOTA	1334		LEU						1.00 20.32
MOTA	· 1335				·	15.558	-10.455	33.272	1.00 17.54
MOTA	1336	CD2	LEU	171		•	-12.464		
ATOM	1337	C	LEU	171	•	18.032	-7.726	31.600	1.00 13.09
ATOM	1338	0 -	LEU	171		17.877	-7.043	32.629	1.00 12.73
MOTA	1339	N	PR0	172		19.128	-7.608	30.836	1.00 12.83
ATOM	1340	CD	PR0	. 172	٠	19.556	-8.419	29.679	1.00 13.47
ATOM.	1341	CA	PR0	172		20.161	-6.633	31.212	1.00 12.48
ATOM	1342	CB	PR0	172		21.238	-6.839	30.147	1.00 12.98
ATOM	1343	CG	PR0	172	•	21.049	-8.280	29.732	1.00 13.76

MOTA	1344	C	PRO .	172	19.673	-5.187	31.274	1.00 12.70
ATOM	1345	0	PRO -	172	20.249	-4.360	31.993	1.00 12.82
MOTA	1346	N	TYR	173	18.624	-4.894	30.521	1.00 11.48
MOTA	1347	CA	TYR	173	18.062	-3.547	30.452	1.00 11.83
MOTA	1348	CB	TYR	173	17.718	-3.207	29.009	1.00 12.45
ATOM	1349	CG	TYR	173	18.897	-3.324	28.087	1.00 12.78
MOTA	1350	CD1	TYR	173	19.693	-2.222	27.820	1.00 13.91
MOTA	1351	CE1	TYR	173	20.812	-2.319	26.989	1.00 14.46
MOTA	1352	CD2	TYR	173	19.236	-4.546	27.501	1.00 13.62
MOTA	1353	CE2	TYR	173	20.347	-4.657	26.668	1.00 15.04
MOTA	1354	CZ	TYR	173			26.419	1.00 15.41
MOTA	1355		TYR		22.231			1.00 18.27
ATOM	1356		A 100		16.771		of the second	and the control of the state of
MOTA				The second second second	at a large to	and the second second	we way to the second	1.00 11.61
MOTA	1358		VAL	The second secon	16.205		And the second of the second	
MOTA	1359			174	14.927			
MOTA				174	14.149			
				174	14.648		The State of the s	1.00 15.43
MOTA				174		great the second	32.580	A contract the state of the contract of the co
MOTA	1363	C			15.052			
MOTA	1364			174	and the second of the second			and the second of the second o
MOTA	1365			175	14.059			
MOTA	1366		*	175		and the second second		
* *	1367			175				1.00 13.92
ATOM	1368	•	ALA				36.743	
ATOM	1369	0	ALA	175	12.995		36.529	
	1370			176				1.00 11.44
	1371		PRO.	176				1.00 12.40
ATOM	1372							1.00 12.63
	1373			176	15.124		40.097	
ATOM	1374		PRO	176	15.164		39.831	
	1375			•				1.00 13.15
ATOM	1376		PRO	176	12.368		39.919	1.00 13.29
ATOM	1377		GLU	177	11.906	· ·	39.346	1.00 12.90
ATOM	1378		GLU	177	10.525		39.788	1.00 14.16
ATOM	1379	CB	GLU	177	9.798		39.740	
ATOM	1380	CG	GLU	177	9.624		38.419	
ATOM	1381		GLU	177.	10.815	-0.414		1.00 13.91
ATOM	1382		GLU	177	10.624	0.519		1.00 13.00
ATOM	1383		GLU	177	11.914	-0.606		1.00 13.08
. ATOM	1384	C	GLU	177	9.746			1.00 15.90
ATOM	1385	0	GLU	177	8.798	-5.064	39.482	1.00 17.12

FIG.11B-33

MOTA	1386	N	LEU	178	10.129	-4.729	37.726	1.00 16.49
ATOM	1387	CA	LEU	178	9.424	-5.742	36.943	1.00 19.28
MOTA	1388	CB	LEU	178	9.957	-5.804	35.506	1.00 22.15
ATOM	1389	CG	LEU	178	9.454	-6.848	34.501	1.00 24.53
MOTA	1390	CD1	LEU	178	10.036	-8.220	34.827	1.00 25.14
MOTA	1391	CD2	LEU	178	7.927	-6.873	34.518	1.00 25.07
MOTA	1392	С	LEU	178	9.622	-7.096	37.565	1.00 20.37
MOTA	1393	0	LEU	178	8.739	-7.954	37.516	1.00 20.98
MOTA	1394	N	LEU	179	10.791	-7.302	38.155	
MOTA	1395	CA	LEU	179	11.101	-8.584	38.766	1.00 21.63
ATOM	1396	CB	LEU	179	12.617	-8.838	38.700	1.00 21.75
MOTA	1397	CG	LEU	179	13.233	-8.817	37.294	1.00 23.23
MOTA	1398	CD1	LEU	179	14.748	-8.931	37.351	1.00 22.87
MOTA	1399	CD2	LEU	179	12.639	-9.954	36.485	1.00 22.80
MOTA	1400	C	LEU	179	10.628	-8.700	40.202	1.00 22.23
ATOM	1401	0_	LEU	179	10.591	-9.799	40.767	1.00 24.11
ATOM	1402	N	LYS	180	10.230	-7.594	40.810	1.00 21.86
ATOM	1403	, CA	LYS	180	9.827	-7.680	42.212	1.00 22.25
MOTA	1404	CB	LYS	180	10.813	-6.909	43.092	1.00 24.33
MOTA	1405	CG	LYS	180	10.945	-5.385	42.935	1.00 27.64
MOTA	1406	CD	LYS	180	11.950	-4.753	43.967	1.00 30.62
MOTA	1407	CE	LYS	180	13.334	-5.432	43.916	1.00 31.90
MOTA	1408	NZ	LYS	180	14.305	-4.830	44.871	1.00 34.05
MOTA	1409	C	LYS	180	8.454	-7.213	42.594	1.00 21.51
MOTA	1410	0	LYS	180	8.007	-7.463		1.00 21.61
MOTA	1411	N	ARG	181	7.760	-6.547	41.680	1.00 20.15
MOTA	1412	CA	ARG	181	6.438	-6.015	•	1.00 19.69
MOTA	1413	CB	ARG	181	6.455	-4.483		1.00 20.79
MOTA	1414	CG	ARG	181	7.705	-3.742	42.554	1.00 23.16
MOTA	1415		ARG	181	8.028	-2.949		
MOTA	1416	NE	ARG	181	7.696	-3.723		1.00 26.61
ATOM	1417	CZ	ARG	181	8.122		46.281	1.00 27.46
MOTA	1418		ARG	181	7.708			1.00 25.45
MOTA	1419	NH2	ARG	181	8.959	•	46.570	1.00 29.25
MOTA	1420	С	ARG	181	5.384		40.995	1.00 19.43
MOTA	1421	0	ARG	•	5.679	-6.774	39.818	1.00 18.33
MOTA	1422	N	ARG	182	,,,,,,	-6.673	41.468	1.00 18.70
MOTA	1423	CA	ARG	182	3.090	-7.125	40.576	1.00 19.47
MOTA	1424	CB	ARG	182	1.813	-7.460	41.348	1.00 22.25
MOTA	1425	CG	ARG	182 ·	0.886	-8.101	40.297	1.00 26.02
MOTA	1426	CD	ARG	182	-0.443	-8.656	40.836	1.00 27.77
MOTA	1427	NE	ARG	182	-1.305	-7.590	41.330	1.00 31.09

MOTA	1428	CZ	ARG	182		-2.507	-7.787	41.859	1.00 33.26
ATOM.	1429	NH1	ARG	182		-2.995	-9.017	41.970	1.00 34.85
ATON	1430	NH2	ARG	182		-3.225	-6.749	42.269	1.00 34.56
MOTA	1431	C .	ARG	182		2.728	-6.068	39.537	1.00 18.40
MOTA	1432	.0	ARG	182		2.482	-6.397	38.372	1.00 19.29
ATOM	1433	: N	GLU	183		2.668	-4.808	39.958	1.00 17.24
MOTA	1434	CA	GLU :	183		2.337	-3.715	39.049	1.00 16.19
MOTA	1435	CB	GLU	183		0.974	-3.102	39.394	1.00 17.54
MOTA	1436	CG	GLU	183		-0.225	-4.044	39.253	1.00 19.75
MOTA	1437	CD	GLU	183	: 1	-1.439	-3.182	39.621	1.00 21.38
MOTA	1438	0E1	GLU	183		-1.593	-2.835	40.813	1.00 23.31
MOTA	1439	0E2	GLU	183	-	-2.208	-2.855	38.697	1.00 21.76
MOTA	1440	C	GLU	183	nga il Vant	3.387	-2.621	39.147	1.00 15.05
MOTA	1441	0	GLU	183		4.085	-2.503	40.148	1.00 13.47
ATOM	1442	N	PHE	184	en.	3.480	-1.797	38.111	1.00 14.19
MOTA	1443	CA	PHE	184		4.474	-0.738	* * * * * * * * * * * * * * * * * * * *	
ATOM	1444	CB	PHE	184		5.861	-1.343	and the second second second	
MOTA	1445	CG	PHE	184			-2.409		
MOTA	1446	CD1	PHE	184	$y_{i_1 i_2 i_3}$		-2.079	***	1.00 14.32
ATOM	1447		PHE	184		5.768	-3.752		1.00 13.96
ATOM		CE1		184		5.688	-3.068	34.441	1.00 14.25
ATOM	> 1449	CE2		184		5.637	-4.754	the contract of the contract o	
MOTA	1450	CZ		184		5.595	-4.407		
ATOM	1451		PHE	•	\$ P. S.	4.138	0.318		and the second second second
MOTA	1452		PHE		ing it	3.427		36.120	•
MOTA	1453		HIS			4.631	1.524		
ATOM	1454		HIS			4.442			
ATOM	1455		HIS			4.892	3.934		.,,
MOTA	1456		HIS	185	•	3.947			
MOTA	1457		•	185		4.013		39.527	
MOTA	1458		HIS	185	· ;	2.770	5.065	37.865	1.00 12.62
	1459				** **				1.00 13.67
	1460		HIS	•		2.886		40.005	
ATOM	1461		HIS	185		5.292			
MOTA	1462		HIS	185	•	6.444		35.236	
MOTA	1463		ALA		•	4.723		34.005	
MOTA	1464		ALA			5.444		32.754	
MOTA	1465		ALA		•	*	•		•
MOTA	1466		ALA			6.677			
MOTA	1467		ALA	186		7.739			
MOTA	1468		GLU	187		6.530			
MOTA	1469	CA	GLU	187	•	7.602	5.548	32.564	1.00 10.72

FIG.11B-35

MOTA	1470	CB	GLU	187		7.133	7.070	32.879	1.00 11.96
ATOM .	1471	CG	GLU	187		6.042	7.443	31.817	1.00 13.69
MOTA	1472	CD	GLU	187		5.429	8.758	32.247	1.00 15.11
MOTA	1473	0E1	GLU .	187		5.768	9.825	31.693	1.00 15.93
MOTA	1474	0E2	GLU	187		4.596	8.703	33.175	1.00 16.67
MOTA	1475	C	GLU	187		8.974	5.371	33.186	1.00 10.14
MOTA	1476	0	GLU	187		9.990	5.441	32.487	1.00 10.23
ATOM	1477	N	PRO	188		9.032	5.065	34.490	1.00 10.27
MOTA	1478	CD	PRO.	188		7.972	5.138	35.507	1.00 10.22
MOTA	1479	CÁ	PRO	188		10.346	4.792	35.105	1.00 10.11
MOTA	1480	CB.	PRO	188		10.013	4.656	36.610	1.00 10.75
MOTA	1481	CG	PRO	188		8.762	5.516	36.770	1.00 9.50
ATOM.	1482	C	PR0	188		11.046	3.548	34.514	1.00 10.42
MOTA	1483	0	PRO-	188		12.261	3.450	34.570	1.00 10.08
MOTA	1484	N	VAL	189		10.284	2.601	33.957	1.00 11.14
MOTA	1485	CA	VAL	189		10.893	1.404	33.363	1.00 10.35
MOTA	1486	CB	VAL	189		9.798	0.368	33.002	1.00 10.10
MOTA	1487	CG1	VAL	189		10.406	-0.821	32.238	1.00 10.42
MOTA	1488	CG2		189		9.118	-0.113	34.271	1.00 11.67
MOTA	1489	C	VAL	189		11.706	1.826	32.106	1.00 10.38
ATOM	1490	0	VAL	189		12.848	1.387	31.906	1.00 10.25
MOTA	1491	N	ASP	190		11.117	2.692	31.284	1.00 10.88
MOTA	1492	CA	ASP	190		11.811	3.165	30.102	1.00 10.93
ATOM	1493	CB	ASP:	190		10.873	3.938	29.160	
ATOM	1494	CG	ASP			9.993	3.059	28.286	1.00 12.68
ATOM	1495	OD1		190		10.297	1.881	28.008	1.00 12.41
ATOM	1496	OD2		190		8.958	3.577	27.818	1.00 13.13
ATOM	1497		ASP	190	·	12.991	4.064	30.512	1.00 10.81
MOTA	1498		ASP	190		14.032	4.050	29.855	1.00 10.56
ATOM	1499		VAL	191	٠,	12.850	4.818	31.603	1.00 10.14
MOTA	1500	CA	VAL	191		13.963	5.665	32.039	1.00 9.64
ATOM	1501							_	1.00 9.54
ATOM	1502		VAL	191	• :		7.130		1.00 10.40
ATOM	1503			191		12.573	•	32.808	1.00 10.05
ATOM	1504		VAL	191		15.173		32.422	1.00 9.21
ATOM			VAL	191	•	16.327			
ATOM	1506	N	TRP	192		14.889	3.691		1.00 9.61
ATOM	1507		.TRP	192	٠.	15.935	2.769	33.572	
ATOM	1508		TRP	192		15.321	1.662	34.439	•
ATOM		CG				16.300	0.619	34.873	1.00 10.53
MOTA	1 510		TRP.			16.870	0.465	36.183	1.00 10.09
ATOM	1511	CE2	TRP	192		17.739	-0.646	36.129	1.00 10.53

FIG.11B-36

MOTA	1512	CE3	TRP	192		16.722	1.150	37.398	1.00 11.26
MOTA	1513	CD1	TRP	192	÷	16.834	-0.371	34.105	1.00 10.22
MOTA	1514	NE1	TRP .	192	:	17.695	-1.135	34.852	1.00 11.11
MOTA	1515	CZ2	TRP	192		18.466	-1.091	37.245	1.00 11.96
MOTA	1516	CZ3	TRP.	192		17.442	0.703	38.514	1.00 11.54
ATOM	1517	CH2	TRP	192	.:	18.305	-0.409	38.421	1.00 10.99
MOTA	1518	'.C . '	TRP	192		16.684	2.150	32.394	1.00 9.91
MOTA	1519	0	TRP	192	especial	17.927	2.133	32.389	1.00 9.89
MOTA	1520	N	SER	193	jt v	15.949	1.619	31.412	1.00 9.48
ATOM	1521	CA	SER	193		16.618	1.031	30.253	1.00 9.06
ATOM	1522	CB	SER	193	٠.	15.610	0.363	29.307	1.00 9.01
MOTA	1523	OG	SER	193		14.587		28.916	
MOTA	1524	C	SER	193	— ••••••	17.463			•
MOTA	1525	0	SER	193		18.520	5 July 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1		
MOTA	1526	N	CYS	194		16.999		29.479	1.00 9.70
MOTA	1527		CYS	4.1		17.796		28.847	
ATOM	1528	CB		194		17.061		28.851	
ATOM	1529		CYS	194		15.742			
MOTA	1530	С	CYS			19.151			
MOTA	1531	0	CYS	194		20.178		29.068	
MOTA	1532	N	GLY	195		19.104			1.00 9.58
MOTA	1533		GLY	195		20.307	for the formal and the first of	31.793	the state of the s
ATOM	1534		GLY	195	Mat 3	21.288			
ATOM	1535		GLY		4	22.498			4 4 40 4 4
MOTA	1536		ILE	196		20.762			
ATOM	1537		ILE	196		21.631			
MOTA	1538		ILE			20.823			
ATOM		CG2		196	din t	19.584		· — · - · · · · · · · · · · · · · · · 	
ATOM		CG1			•	21.610			
MOTA		CD1		196	•• • •			31.133	
MOTA	1542		ILE	196		22.249		29.493	1.00 11.39
	-			196				29.249	· · · · · · · · · · · · · · · · · · ·
MOTA	1544			197			· ·	28.592	
MOTA				197		21.989		27.257	
MOTA		CB						26.358	
	1547			197	•	21.465			1.00 10.47
ATOM		CG2				19.911		25.963	
MOTA	1549		VAL			23.129		27.398	1.00 10.49
MOTA	1550		VAL			24.161		26.726	1.00 10.94
MOTA	1551		LEU	198		22.944			
MOTA	1552		LEU	198		23.977			
MOTA	1553	CB	LEU	198		23.537	6.200	29.571	1.00 12.55

FIG.11B-37

•	ATOM	1554	CG	LEU	198	23.879	7.664	29.261	1.00 16.51
	MOTA	1555	CD1	LEU	198	23.869	8.361	30.615	1.00 12.50
	MOTA	1556	CD2	LEU	198	25.154	7.918	28.500	1.00 13.83
	MOTA	1557	С	LEU	198	25.253	4.511	29.047	1.00 11.79
	MOTA	1558	0	LEU	198	26.371	4.801	28.600	1.00 11.82
	MOTA	1559	N	THR .	199	25.067	3.592	29.985	1.00 11.20
	ATOM	1560	CA	THR	199	26.199	2.862	30.574	1.00 11.87
	ATOM	1561	CB	THR	199	25.699	1.860	31.641	1.00 11.32
•	ATOM	1562	0G1	THR :	199	25.041	2.585	32.677	1.00 11.23
	ATOM	1563	CG2	THR	199	26.878	1.088	32.291	1.00 11.31
	MOTA	1564	C.	THR	199	26.947	2.154	29.486	1.00 12.08
	MOTA	1565	0,	THR	19 9	28.181	2.237	29.410	1.00 12.74
	ATOM	1566	N	ALA	200	26.202	1.474	28.614	1.00 13.09
	ATOM	1567		ALA	200	26.805	0.737	27.506	1.00 13.17
	ATOM	1568	CB	ALA .	200	25.720	0.002	26.712	1.00 12.95
	ATOM	1569	C	ALA	200	27.589	1.668	26.568	1.00 14.04
	MOTA	1570	0	ALA	200	28.690	1.345	26.140	1.00 13.51
	MOTA	1571	N	MET	201	27.023	2.822	26.241	1.00 12.86
	ATOM	1572	CA	MET	201	27.725	3.728	25.335	1.00 12.63
	MOTA	1573	CB	MET	201	26.849	4.930	24.954	1.00 12.56
	MOTA	1574	CG	MET	201	25.592	4.544	24.110	1.00 13.61
	MOTA	1575	SD	MET	201	24.831	6.026	23.390	1.00 12.69
	MOTA	1576	CE	MET	201	24.080	6.854	24.850	1.00 12.19
	MOTA		C	MET	201	29.011	4.268		1.00 12.12
	MOTA	1578	0	MET	201	29.997	4.484	25.222	1.00 12.45
	MOTA	1579	N	LEU	202	29.019	4.458	27.247	1.00 11.16
	MOTA	1580	CA	LEU	202	30.199	5.014	27.907	1.00 12.42
	MOTA	1581	CB	LEU	202	29.782	5.864		1.00 12.19
	MOTA	1582	CG	LEU	202	28.994	7.113	28.691	1.00 11.96
	ATOM	1583		LEU	202	28.551	7.912	29.931	1.00 13.07
	MOTA	1584		LEU	202	29.891	7.960	27.796	1.00 12.51
	MOTA	1585	C	LEU	202	31.262		28.384	•
	ATOM	1586	. 0	LEU	202	32.414	4.384	28.610	1.00 14.80
	ATOM	1587		ALA	203		2.734	28.514	1.00 13.10
	ATOM	1588	CA	ALA	203	31.839	1.726	28.988	1.00 15.17
	ATOM	1589	CB	ALA	203	31.424	1.252	30.367	1.00 14.79
	MOTA	1590	C	ALA	203	32.004	0.521	28.047	1.00 15.81
	MOTA	1591	0	ALA	203	32.926	-0.290	28.216	1.00 17.03
	MOTA	1592	N	GLY	204	31.117	0.394	27.070	1.00 15.77
	ATOM	1593	CA	GLY	204	31.218	-0.728	26.149	1.00 17.71
	ATOM .	1594	. C	GLY	204	30.957	-2.072	26.803	1.00 18.37
	ATOM	1595	0	GLY	204	31.451	-3.112	26.340	1.00 19.10

MOTA	1596	N GLU	205	30.199	-2.052	27.888	1.00 18.22
MOTA	1597	CA GLU	205	29.850	-3.268	28.610	1.00 19.72
MOTA	1598	CB GLU	205	30.977	-3.692	29.552	1.00 22.13
ATOM	1599	CG GLU	205	31.134	-3.004	30.896	1.00 24.83
MOTA	1600	CD GLU	205	32.225	-3.740	31.729	1.00 25.92
MOTA	1601	OE1 GLU	205	32.102	-4.890	32.202	1.00 28.12
MOTA	1602	OE2 GLU	205	33.274	-3.121	31.912	1.00 26.08
MOTA	1603	C GLU	205	28.582	-3.039	29.424	1.00 18.53
MOTA	1604	O GLU	205	28.292	-1.915	29.845	1.00 18.22
MOTA	1605	N LEU	206	27.819	-4.107	29.622	1.00 17.56
ATOM	1606	CA LEU	206	26.579	-4.045	30.396	1.00 17.14
MOTA	1607	CB LEU	206	25.563	-5.054	29.847	1.00 17.00
ATOM	1608	CG LEU	206	25.030	-4.728	28.447	1.00 18.03
ATOM	1609	CD1 LEU	206	26.152	-4.419	27.471	1.00 22.13
MOTA	1610	CD2 LEU	206	24.233	-5.927	27.948	1.00 17.79
MOTA	1611	C LEU	206	26.976	-4.351	31.811	1.00 16.84
ATOM	1612	O LEU	206	27.782		32.057	1.00 16.85
ATOM	1613	N PRO	207	26.420	-3.604		1.00 16.25
MOTA	1614	CD PRO	207	25.415			
MOTA	1615	CA PRO	207	26.712	-3.757		
MOTA	1616	CB PRO	207	26.077	-2.503		
MOTA	1617	CG PRO	207	24.870	-2.295		The state of the s
MOTA	1618	C PRO	207	26.305	-5.042	34.871	1.00 17.01
MOTA	1619	O PRO	207	27.012	-5.518	الوديد أن والحداث	10.00
MOTA	1620	N TRP	208	25.181	-5.626		1.00 16.29
MOTA	1621	CA TRP	208	24.726		35.074	
MOTA	1622		208	24.006	-6.564		
MOTA	1623	CG TRP	.208	23.028	-5.406	36.304	1.00 15.34
MOTA	1624	CD2 TRP	208	23.198	•		
MOTA	1625	CE2 TRP	208		-3.323		
ATOM	1626	CE3 TRP	208	24.186	-3.530		
		CD1 TRP					1.00 15.76
MOTA	1628		208	21.273			
			208				
MOTA		•	208	24.037	•	•	• •
MOTA		CH2 TRP	208	•		• • •	
MOTA	1632		·			,	1.00 17.33
MOTA	1633		208				
ATOM		N ASP	209		-9.013		•
MOTA	•	· CA · ASP	209	22.885			
ATOM	1636		209				1.00 22.30
MOTA	1637	· CG ASP	209	24.466	-11.974	-34.039	1.00 23.93

FIG.11B-39

	ATOM	1638	OD1	ASP	209	25.057 -11.638	32.994	1.00 25.16
٠,	MOTA	1639	OD2	ASP	209	24.939 -12.771	34.884	1.00 27.15
	MOTA	1640	C	ASP	209	21.406 -9.587	33.984	1.00 18.77
	ATOM	1641	0	ASP	209	20.604 -9.675	33.052	1.00 17.74
	MOTA	1642	N	GLN	210	21.068 -9.178	35.205	1.00 18.55
	MOTA	1643	CA	GLN	210	19.712 -8.775	35.559	1.00 19.19
	MOTA	1644	CB	GLN	210	18.805 -10.003	35.664	1.00 20.25
	ATOM	1645	CG	GLN	210	19.377 -11.006	36.658	1.00 21.64
	ATOM	1646	CD	GLN	210	18.489 -12.229		1.00 23.03
	MOTA	1647	0E1	GLN	210	18.452 -12.918	35.555	1.00 23.31
	MOTA	1648	NE2	GLN	210	17.765 -12.503	37.650	1.00 24.19
	ATOM	1649	C	GLN	210	19.775 -8.010	36.865	1.00 19.59
	MOTA	1650	0	GLN	210	20.691 -8.209	37.669	1.00.18.71
	MOTA	1651	N	PRO	211	18.806 -7.111	37.105	1.00 18.57
	MOTA	1652	CD.	PRO	211	17.799 -6.619	36.150	1.00 18.07
	MOTA	1653	CA	PRO:	211	18.783 -6.311	38.334	1.00 19.80
٠	MOTA	1654	CB	PRO	211	17.999 -5.071	37.895	1.00 19.10
	MOTA	1655	CG	PRO	211	16.995 -5.644	37.004	1.00 18.49
	MOTA	1656	С	PRO	211	18.202 -7.039	39.533	1.00 21.25
	MOTA	1657	0	PRO	211	17.149 -6.664	40.049	1.00 20.53
	ATOM	1658	N	SER	212	18.914 -8.068	39.986	1.00 22.92
	MOTA	1659	CA	SER	212	18.476 -8.876	41.122	1.00 25.26
	ATOM	1660	CB	SER	212	18.232 -10.322	40.696	1.00 25.71
	MOTA	1661	-OG	SER	212	17.269 -10.404	39.656	1.00 27.04
	MOTA	1662	C	SER	212	19.540 -8.909	42.200	1.00 26.27
	ATOM	1663	· 0 ·	SER	212	20.728 -8.823	41.911	1.00 26.27
	ATOM	1664	N	ASP	213	19.112 -9.031	43.449	1.00 28.30
	ATOM	1665	CA	ASP	213	20.060 -9.091	44.558	1.00 30.12
	MOTA	1666	CB	ASP	213	19.308 -9.155	45.885	1.00 31.41
	MOTA	1667	CG	ASP	213	18.700 -7.785	46.123	1.00 32.87
	ATOM	1668	0D1	ASP	213	17.794 -7.695	46.971	1.00 34.35
	MOTA	1669	0D2	ASP	213	19.131 -6.806		1.00 33.85
	MOTA	1670	C	ASP	213	20.950 -10.325	44.402	1.00 30.47
	MOTA	1671	0	ASP	213	22.085 -10.347	44.881	1.00 30.66
	MOTA	1672	N	SER	214	20.431 -11.345	43.722	1.00 30.65
	ATOM	1673	CA	SER	214	21.182 -12.577		1.00 30.71
	MOTA	1674	CB	SER	214	20.266 -13.676		1.00 31.27
	ATOM	1675	OG	SER	214	19.713 -13.315		1.00 33.24
	MOTA	1676	C	SER	214	22.295 -12.353		1.00 30.15
	MOTA	1677	0	•	- 214	23.180 -13.201		1.00 30.59
	MOTA	1678	N	CYS	215	22.237 -11.212		1.00 28.37
	ATOM	1679	CA	CYS	215	23.248 -10.857		1.00 27.11
						•		

FIG.11B-40

	MOTA	1680	CB	CYS	215	22.615	-10.052	39.679	1.00 27.35
	MOTA	1681	SG	CYS	215	23.795	-9.692	38.381	1.00 24.98
	MOTA	1682	C	CYS	215	24.290	-10.029	41.524	1.00 26.85
	ATOM	1683	0	CYS	215	24.046	-8.879	41.881	1.00 25.26
	MOTA	1684	N	GLN .	216	25.465	-10.618	41.730	1.00 26.11
	ATOM	1685	CA	GLN	216	26.547	-9.945	42.432	1.00 25.73
	ATOM	1686	CB	GLN	216	27.806	-10.824	42.396	1.00 27.39
	ATOM	1687	CG	GLN		28.908	-10.267	43.303	1.00 29.33
		1688	CD	GLN	216	28.445	-10.161	44.773	1.00 30.23
	MOTA	1689	0E1	GLN	216	28.777	-9.201	45.469	1.00 31.39
	ATOM	1690	NE2	GLN	216	27.691	-11.153	45.236	1.00 29.72
	ATOM	1691	C	GLN	.216	26.867	-8.526	41.914	1.00 24.91
	ATOM	1692	0	GLN	216	27.064	-7.606	42.705	1.00 24.59
	ATOM	1693	N	GLU	217	26.904			and the second s
١.	MOTA	1694	CA	GLU	217	6 4 14 14			1.00 22.82
	MOTA	1695	CB	GLU	217			38.490	1.00 23.60
	MOTA	1696	CG	GLU -	217	28.545	-7.945	37.893	
Ì	ATOM	1697	CD	GLU	217		-9.382		1.00 26.01
	ATOM				217				1.00 26.36
	ATOM	1699	0E2	GLU	217				1.00 27.79
	ATOM	1700			217			40.387	
	MOTA	1701	0 3	GLU	217			40.597	all the contract of the contra
÷	MOTA	1702		TYR				40.467	
	ATOM	1703	CA	TYR	218				1.00 21.45
		1704		•	218				1.00 20.94
•	MOTA	1705		TYR	218	. "			1.00 20.26
		1706		TYR		21.289		40.185	7 × × × × × ×
	ATOM			LTYR				with the state of the state of	
					218				e de la companya de l
	ATOM	• •			218			42.081	
	MOTA	1710		•	218				1.00 19.91
		1711			218			•	1.00 20.57
	MOTA	1712		TYR	218			42.332	1.00 22.12
	MOTA	1713			218	23.853		42.706	1.00 21.46
	ATOM	1714			219	24.26]		43.178	1.00 22.64
	MOTA	1715			219	24.423		44.605	1.00 23.55
	ATOM	1716		SER				45.403	1.00 24.17
	. Atom	1717		SER	219	23.642			1.00 26.99
	ATOM	1718		SER	219	25.580			1.00 23.48
	ATOM	1719		SER	219	25.483		45.578	1.00 24.10
	ATOM	1720		ASP	220				
	ATOM	1721	CA	ASP	220	27.837	7 -4.240	44.169	1.00 22.96

FIG.11B-41

	ATOM	1722		ASP	220	28.941	-4.732	43.232	1.00 24.39
	ATOM	1723	CG	ASP -	220	29.580	-5.983	43.835	1.00 25.92
	MOTA	1724	OD1	ASP	220	30.398	-6.603	43.128	1.00 28.41
	MOTA	1725	002	ASP -	220	29.278	-6.340	44.992	1.00 27.22
	ATOM	1726	С	ASP	220	27.480	-2.786	43.828	1.00 22.40
	ATOM	1727	0	ASP	220	28.005	-1.855	44.428	1.00 22.30
	MOTA	1728	N.	TRP	221	26.585	-2.594	42.865	1.00 21.17
	MOTA	1729	CA	TRP	221	26.179	-1.241	42.498	1.00 20.70
	MOTA	1730	CB ,	TRP	221	25.391	-1.291	41.176	1.00 19.27
	ATOM	1731	CG	TRP	221	24.638	-0.020	40.833	1.00 17.73
	MOTA	1732	CD2	TRP	221	25.191	1.230	40.395	1.00 16.84
	ATOM	1733	CE2	TRP .	221	24.106	2.117	40.187	1.00 17.43
	MOTA	1734	CE3	TRP	221	26.491	1.688	40.154	1.00 17.01
	MOTA	1735	CD1	TRP	221	23.287	0.156	40.874	1.00 17.87
	MOTA	1736	NE1	TRP	221	22.959	1.435	40.491	1.00 17.27
	ATOM	1737	CZ2	TRP	221	24.284	3.438	39.747	1.00 16.90
	MOTA	1738	CZ3	TRP	221	26.668	3.013	39.715	1.00 16.65
	MOTA	1739	CH2	TRP	221	25.573	3.864	39.518	1.00 17.11
	MOTA	1740	C	TRP	221	25.376	-0.599	43.651	1.00 21.75
	MOTA	1741	0	TRP	221	25.617	0.552	44.015	1.00 21.20
	MOTA	1742	N	LYS	222	24.441	-1.351	44.225	1.00 23.83
	MOTA	1743	CA	LYS	222	23.641	-0.828	45.324	1.00 26.15
	MOTA	1744	CB	LYS	222	22.564	-1.831	45.735	1.00 26.74
	ATOM	1745	∵CG.	LYS	222	21.471	-1.821	44.636	1.00 27.05
	ATOM	1746	CD	LYS	222	20.119	-2.467	45.022	1.00 28.62
	ATOM	1747	CE	LYS	222	20.199	-3.943	45.413	1.00 28.17
	MOTA	1748	NZ	LYS	222	18.869	-4.443	45.862	1.00 30.18
	MOTA	1749	C	LYS	222	24.524	-0.497	46.528	1.00 27.58
•	ATOM	1750	0	LYS	222	24.150	0.320	47.371	1.00 27.87
	MOTA	1751	N	GLU	223	25.694	-1.126	46.586	1.00 29.19
	MOTA	1752	CA	GLU	223	26.650	-0.902	47.670	1.00 30.71
	ATOM	1753	CB"	GLU	223	27.426	-2.187	47.975	1.00 32.35
	MOTA	1754	CG.		223				1.00 35.04
	MOTA	1755	CD	GLU	223	27.341	-4.629	48.610	1.00 36.39
•	ATOM	1756	:0E1	GLU	223	28.026	-4.647		1.00 37.89
	ATOM	1757	OE2	GLU .		27.315		47.799	
	ATOM	1758	C	GLU					1.00 31.11
	ATOM	1759	0	GLU	223	28.595	0.476		1.00 31,11
	ATOM	1760		LYS	224	27.414	0.835		1.00 31.36
	ATOM	1761	CA	LYS	224	28.250	1.935		1.00 31.60
	MOTA	1762	CB	LYS	224		3.084		1.00 32.70
	MOTA	1763	CG	LYS	224	26.902			1.00 34.17

FIG.11B-42

					and the second second sections with			Contraction with the second of
MOTA	1764	CD	LYS -	224	25.731	2.967	47.318	1.00 35.15
MOTA	1765	CE	LYS	224	25.845	2.601	48.823	1.00 36.31
ATOM	1766	NZ	LYS	224	25.781	3.822	49.677	1.00 37.18
MOTA	1767	C	LYS	224	29.720	1.607	45.343	1.00 31.34
MOTA	1768	0	LYS	224	30.595	2.467	45.463	1.00 31.10
MOTA	1769	N	LYS	225	29.982	0.377	44.918	1.00 30.93
MOTA	1770	CA	LYS	225	31.347	-0.028	44.574	1.00 31.37
MOTA	1771	CB	LYS	225	31.493	-1.543	44.742	1.00 31.69
MOTA	1772	CG	LYS	225	31.227	-1.904	46.232	1.00 32.87
MOTA	1773	CD	LYS	225	31.162	-3.409	46.591	1.00 33.45
MOTA	1774	CE	LYS	225	32.345	-4.339		1.00 35.03
ATOM	1775	NZ	LYS	225	32.064	-5.731		1.00 36.16
MOTA	1776	C	LYS	225	31.641	24 C		1.00.30.98
MOTA	1777	0	LYS	225	31.751	-0.465		1.00 30.71
MOTA	1778	N	THR	226	31.766		42.886	1.00 31.21
MOTA	1779	CA	THR	226	32.009	2.208		1.00 31.56
MOTA	1780	CB	THR	226	31.458			1.00 31.65
MOTA	1781		THR		31.977		42.478	1.00 32.08
MOTA	1782	CG2	THR	226	29.939		41.514	1.00 31.20
MOTA	1783	C	THR	226	33.464		41.108	
MOTA	1784	0	THR	226	33.869	2.803		
ATOM	1785	N	TYR	227	34.252			1.00 32.46
MOTA	1786		TYR	227	35.653		41.456	1.00 32.59
MOTA	1787		TYR	227	36.518	1.115	and the second transfer	1.00 33.51
MOTA	1788	CG	TYR	227	36.011	•	43.801	1.00 34.46
ATOM	1789		. TYR	227		-1.196	43.703	
MOTA	1790		TYR	227	35.714		44.699	1.00 35.37
ATOM	1791		TYR	227	35. <u>351</u>	a to the same of the same of	44.920	
MOTA	1792			227	34.874	-0.154		1.00 35.15
MOTA	1793		TYR		35.058			
ATOM	1794		TYR	227	34.576			1.00 35.61
ATOM	1795				and the second second			1.00 32.78
MOTA	1796		TYR	227	•		40.254	
,	1797		LEU	228	34.614		40.464	
MOTA	1798				34.517		39.709	
ATOM	1799			228	33.390			•
ATOM	1800			228			41.675	
ATOM	1801		1 LEU	. 228	32.676			
ATOM			2 LEU	228	35.116			
ATOM	1803		LEU	228	34.264			
MOTA	1804		LEU	228	33.627			
MOTA	1805	N	ASN	229	34.762	2.640	37.387	1.00 35.26

FIG.11B-43

ATOM	1806	CA	ASN	229	34.716	-2.588	35.925	1.00 35.58
MOTA	1807	CB	ASN	229	34.458	-3.987	35.346	1.00 36.34
MOTA	1808	CG	ASN	229	35.512	-4.243	34.249	1.00 37.55
ATOM .	1809	OD1	ASN	229	36.703	-4.002	34.455	1.00 37.91
MOTA	1810	ND2	ASN	229	35.069	-4:742	33.096	1.00 37.37
ATOM	1811	C	ASN	229	33.829	-1.633	35.198	1.00 34.28
MOTA	1812	0.	ASN	229	34.300	-0.626	34.665	1.00 35.89
ATOM	1813	N	PRO	230	32.516	-1.897	35.159	1.00 33.61
MOTA	1814	CD	PRO	230	31.668	-2.722	36.038	1.00 32.77
MOTA	1815	CA	PRO	230	31.718	-0.924	34.408	1.00 30.57
MOTA	1816	CB	PRO	230	30.287	-1.447	34.623	1.00 32.30
MOTA	1817	CG	PRO	230	30.340	-1.971	36.006	1.00 33.38
MOTA	1818	C	PRO	230	31.960	0.575	34.781:	1.00 27.81
MOTA	1819	0	PRO	230	32.499	1.367	33.990	1.00 26.92
MOTA	1820	N	TRP	231	31.578	0.918	35.999	1.00 24.76
MOTA	1821	CA	TRP	231	31.652	2.276	36.514	1.00 22.97
MOTA	1822	CB	TRP	231	30.995	2.295	37.899	1.00 21.67
MOTA	1823	CG	TRP	231	29.833	1.331	37.961	1.00 19.36
MOTA	1824	CD2	TRP	231	28.622	1.407	37.204	1.00 18.86
MOTA	1825	CE2	TRP	231	27.878	0.239	37.485	1.00 18.66
MOTA	1826	CE3	TRP	231	28.095	2.350	36.310	1.00 17.74
MOTA	1827	CD1	TRP	231	29.773	0.155	38.660	1.00 19.31
ATOM	1828		TRP	231	28.605	-0.509	38.377	1.00 17.81
MOTA	1829			. 231	26.634	-0.012	36.904	1.00 17.32
MOTA	1830		TRP	231	26.856	2.103	35.727	1.00 18.02
MOTA	1831		TRP	231	26.139	0.928	36.031	1.00 17.78
ATOM	1832	C	TRP	231	33.033	2.946	36.558	1.00 22.63
ATOM		. 0	TRP	231	33.128	4.159		1.00 22.81
MOTA	1834	N	LYS	232	34.091	2.170	36.754	1.00 22.60
MOTA	1835	CA	LYS	232	35.428	2.764	36.826	1.00 22.75
MOTA	1836		LYS	232	36.477	1.704	37.189	1.00 24.11
ATOM .			LYS	232			36.346	
ATOM	1838		LYS	232		-0.509	37.036	1.00 27.06
MOTA	1839		LYS	232	37.996		36.213	1.00 27.57
ATOM	1840	NZ	LYS	232	39.033	-2.612	36.876	1.00 28.69
MOTA	1841	C	LYS	232	35.860	3.438	35.529	1.00 22.63
MOTA	1842	0	LYS	232	36.790	4.258	35.522	1.00 22.71
MOTA	1843	N	LYS	233	35.182	3.104	34.435	1.00 20.74
- ATOM	1844	CA	LYS	233	35.522	3.658	33.122	1.00 20.06
MOTA	1845	CB	LYS	233	35.264	2.634	32.011	1.00 21.41
MOTA	1846	CG	LYS	233	36.100			1.00 22.20
ATOM	1 847	CD	LYS	233	35.874	0.176	31.106	1.00 23.44

FIG.11B-44

MOTA	1848	CE	LYS	233	34.596	-0.617	31.187	1.00 24.11
MOTA	1849	NZ	LYS	233	34.558	-1.720	30.177	1.00 24.50
MOTA	1850	C	LYS	233	34.855	5.008	32.889	1.00 19.90
ATOM	1851	0	LYS	233	35.272	5.662	31.949	1.00 18.76
MOTA	1852	N	ILE.	234	33.741	5.339	33.521	1.00 20.48
MOTA	1853	CA	ILE	234	32.935	6.424	32.978	1.00 21.37
ATOM	1854	CB	ILE	234	31.491	6.107	33.470	1.00 20.52
ATOM	1855		ILE	234	30.511	7.217	33.130	1.00 20.33
ATOM	1856		ILE	234	31.126	4.755	32.836	1.00 19.24
MOTA	1857	CD1		234	29.665	4.267	33.021	1.00 18.12
MOTA	1858	C	ILE	234	33.591	7.883	33.073	1.00 23.34
ATOM	1859	0	ILE	234	34.414	8.250	32.223	1.00 26.22
MOTA	1860	N	ASP	235	33.190	8.659	34.058	1.00 23.93
ATOM	1861	CA	ASP	235	33.700	10.001	34.367	1.00 21.53
MOTA	1862	CB	ASP	235	33.670	10.978	33.182	1.00 23.07
MOTA	1863	CG	ASP	235	34.063	12.339	33.827	1.00 24.23
MOTA	1864	OD1	ASP	235	33.209	13.262	33.921	1.00 23.91
ATOM	1865		ASP	235	35.237	12.473	34.266	1.00 23.71
MOTA	1866	C	ASP	235	32.742	10.372	35.366	1.00 21.02
ATOM	1867	0	ASP	235	31.577	10.002	35.253	1.00 18.48
ATOM	1868	N	SER	236	33.180	11.101	36.387	1.00 20.18
ATOM	1869	CA	SER	236	32.301	11.481	37.480	1.00 20.49
ATOM	1870	CB	SER	236	33.036	12.390	38.481	1.00 21.41
MOTA	1871	OG	SER	236	33.526	13.563	37.863	1.00 23.42
MOTA	1872	C	SER	236	30.995	12.139	37.117	1.00 18.82
MOTA	1873	0	SER	236	29.971	11.832	37.730	
MOTA	1874	N	ALA	237	31.019		36.129	
MOTA	1875	CA	_ALA_	237	29.825		35.701	
MOTA	1876	CB	ALA.	237	30.194			1.00 16.94
MOTA	1877	C	ALA	237	28.709		the state of the s	1.00 15.31
MOTA	1878	0	ALA	237				1.00 15.03
MOTA	1879	N	PRO	238	·			1.00 14.33
MOTA	1880	CD	PRO	238	30.153			1.00 13.45
MOTA	1881	CA		238	27.908			
MOTA	1882	CB	PRO	238	28.424		32.335	
MOTA	1883		PRO	238	29.934			
MOTA	1884		PRO	. 238	27.584		34.714	
MOTA -	1885		PRO	238	26.461		34.799	•
MOTA	1886		LEU	239	28.579		35.509	
MOTA	1887			239	28.363			
MOTA	1888			239	29.702			
MOTA	1889	CG	LEU	239 -	29.797	7.069	38.059	1.00 17.03

FIG.11B-45

MOTA	1890	CD1 LEU	239		29.461	7.543	39.426	1.00 19.97
ATOM	1891	CD2 LEU	239		28.941	5.890	37.632	1.00 16.99
MOTA	1892	C LEU	239		27.350	9.209	37.548	1.00 14.32
MOTA	1893	O LEU	239		26.521	8.451	38.053	1.00 13.88
MOTA	1894	N ALA	240		27.410	10.513	37.836	1.00 13.60
MOTA	1895	CA ALA	240		26.474	11.121	38.778	1.00 14.03
MOTA	1896	CB ALA	240		26.834	12.596		1.00 14.41
MOTA	1897	C ALA	240	: .	25.049	11.017	38.214	1.00 13.80
MOTA	1898	O ALA	240	: .	24.105	10.815	38.959	1.00 15.01
MOTA	1899	N LEU	241		24.911	11.141	36.898	1.00 13.56
MOTA	1900	CA LEU	241		23.586	11.029	36.289	1.00 12.69
MOTA	1901	CB LEU	241		23.612	11.492	34.824	1.00 12.29
MOTA	1902	CG LEU	241		22.307		34.050	1.00 12.22
MOTA	1903	CD1 LEU	241		21.101	11.892	34.673	1.00 13.19
MOTA	1904	CD2 LEU	241		22.521	11.700	32.616	1.00 11.55
MOTA	1905	C LEU	241		23.144		36.389	1.00 12.74
MOTA	1906	O LEU	241		21.992	9.298	36.744	1.00 13.22
MOTA	1907	N LEU	242		24.051	8.658	36.086	1.00 12.48
ATOM	1908	CA LEU	242		23.697	7.242	36.165	1.00 13.46
MOTA	1909	CB LEU	242		24.901	6.395	35.710	1.00 14.09
MOTA	1910	CG LEU	242		24.951	5.527	34.437	1.00 17.22
MOTA	1911	CD1 LEU	242		23.861	5.790	33.451	1.00 13.60
MOTA	1912	CD2 LEU	242		26.335	5.691	33.831	1.00 14.06
MOTA	1913	C LEU	242	•	23.256	6.879	37.622	
MOTA	1914	O LEU	242		22.369	6.034	37.834	1.00 13.32
MOTA	1915	N HIS	243	٠	23.861	7.526	38.615	1.00 14.11
MOTA	1916	CA HIS	243	:	23.485	7.251	40.004	1.00 14.54
MOTA	1917	CB HIS	243	. :	24.385	8.001	40.998	1.00 16.02
MOTA	1918	CG HIS	243		25.597	7.228	41.426	1.00 18.57
ATOM	1919	CD2 HIS	243		26.911	7.424	41.173	1.00 20.62
MOTA	1920	ND1 HIS	243	• • • •	25.524	6.099	42.216	1.00 20.36
ATOM	1921	CE1 HIS	_ 10		26.743	5.632	42.427	
ATOM	1922	NE2 HIS	243	*	27.603	6.418	41.804	1.00 20.07
ATOM	1923	C HIS	243		22.037	7.679	40.279	1.00 14.42
ATOM	1924	O HIS	243		21.400	7.148	41.181	1.00 15.58
ATOM	1925	N LYS	244		21.548	8.652	39.513	1.00 13.04
ATOM	1926	CA LYS	244	•	20.181	9.138	39.662	1.00 12.66
ATOM	1927	CB LYS	244		20.061	10.586	39.215	1.00 13.26
ATOM	1928	CG LYS	244		20.819	11.500	40.245	1.00 13.78
ATOM	1929	CD LYS	244		20.709	12.997	39.911	1.00 16.39
ATOM	1930	CE LYS	244		21.462	13.451		1.00 16.70
ATOM	1931	NZ LYS	244		21.476	14.962	38.449	1.00 17.30

FIG.11B-46

MOTA	1932	C	LYS	244	19.164	8.319	38.858	1.00 12.13
MOTA	1933	0	LYS	244	17.993	8.241	39.224	1.00 12.40
MOTA	1934	N	ILE :	245	19.621	7.723	37.759	1.00 12.05
MOTA	1935	CA	ILE	245	18.747	6.906	36.912	1.00 10.91
MOTA	1936	CB	ILE	245	19.281	6.819	35.449	1.00 10.95
ATOM		CG2		245	18.405	•	34.617	1.00 10.91
ATOM	1938		ILE		19.237	8.197	34.786	1.00 11.13
ATON	1939		ILE :		19.900		33.354	1.00 12.03
MOTA	1940	С		245	18.606	5.451	37.419	1.00 12.05
ATOM		0		245	17.502		37.489	1.00 13.24
MOTA	1942				19.726	4.831	37.777	1.00 12.15
ATOM		CA		246				1.00 12.12
MOTA	1944			246		2.757.	37.761	1.00 11.77
ATOM				246				1.00 11.06
ATOM	1946		LEU					1.00 10.74
ATOM	1947					2.158	35.515	1.00 11.92
ATOM	1948	C	LEU	246	19.442	3.277	39.711	1.00 13.39
MOTA	1949	0	LEU	246	20.212	2.652	40.468	1.00 14.42
MOTA	1950	N	VAL	247	18.321			1.00 12.95
ATOM	1951	CA	VAL	247			41.483	
ATOM	1952	CB	VAL	247				1.00 14.69
MOTA	1953	CG1	VAL	247	16.472			
MOTA	1954	CG2	VAL	247				
MOTA	1955	C	VAL	247	16.780			
MOTA	1956	0	VAL	247				1.00 13.95
MOTA				248		* .,	42.468	
MOTA	1958	CA	GLU	248	16.008			1.00 15.38
MOTA	1959	CB	GLU	248			43.723	The state of the state of
MOTA	1960	CG	GLU	248			43.749	
MOTA	1961	CD	GLU	248			44.545	
MOTA	1962	0E1	GLU	248	16.574			1.00 27.75
MOTA	1963	0E2	GLU	248			***	1.00 27.17
MOTA	1964	C	GLU	248	•			1.00 14.82
MOTA	1965	0	GLU	248	13.649			1.00 13.94
MOTA	1966	N		249	14.246		43.663	
ATOM	1967	CA		249	12.878			1.00 14.73
ATOM	1968		ASN	249	12.821			1.00 15.18
MOTA	1969		ASN	249	11.435	3.822		
ATOM	1970		L ASN	249	10.499	3.664		1.00 16.50
MOTA	1971		2 ASN	249	11.312		46.643	
ATOM	1972	C	ASN		12.447		42.759	
ATOM	1973	0	ASN	249	12.961	4.438	42.611	1.00 14.63

FIG.11B-47

		• •						
MOTA	1974	N	PRO	250	11.518	2.862	41.912	1.00 13.20
MOTA	1975	CD	PRO	250	10.763	1.599	42.012	1.00 12.72
MOTA	1976	CA	PRO	250	11.057	3.658	40.766	1.00 13.23
MOTA	1977	CB	PRO	250	10.079	2.706	40.055	1.00 13.24
MOTA	1978	CG	PRO	250	9.507	1.906	41.190	1.00 13.59
MOTA	1979	C	PRO.	250	10.446	4.976	41.155	1.00 13.43
MOTA	1980	0	PR0	250	10.442	5.921	40.365	1.00 13.46
MOTA	1981	N	SER	251	9.904	5.050	42.368	1.00 14.06
MOTA	1982	CA	SER	251	9.303	6.302	42.803	1.00 15.63
MOTA	1983	CB	SER	- 251	8.386	6.059	44.002	1.00 15.27
MOTA	1984	OG	SER	251	7.238	5.337	43.589	1.00 15.42
MOTA	1985	C	SER	251	. 10.372	7.369	43.132	1.00 15.99
ATOM	1986	0	SER	251	10.099	8.558	43.044	1.00 19.15
MOTA	1987	N	ALA	252	11.577	6.933	43.480	1.00 16.36
MOTA	1988	CA	ALA	252	12.670	7.846	43.812	1.00 15.52
ATOM	1989	CB	ALA	252	13.504	7.261	44.950	1.00 15.97
MOTA	1990	C .	ALA	252	13.568	8.099	42.602	1.00 14.86
MOTA	1991	0	ALA	252	14.398	9.002	42.603	1.00 16.55
MOTA	1992	N '	ARG	253	13.407	7.279	41.577	1.00 14.26
MOTA	1993	CA	ARG	253	14.230	7.395	40.364	1.00 13.52
MOTA	1994	CB	ARG	253	13.892	6.245	39.416	1.00 13.08
MOTA	1995	CG	A RG	253	14.732	6.070	38.114	1.00 13.50
MOTA	1996	CD	ARG	253	14.277	4.765	37.436	1.00 12.90
ATOM	1997	NE	ARG	253	14.298	3.661	38.395	1.00 13.33
MOTA	1998	CZ	ARG	253	13.564	2.561	38.289	1.00 13.77
MOTA	1999		ARG	253	13.638	1.625	39.238	1.00 12.46
MOTA	2000	NH2	ARG	253 .	12.771	2.397	37.234	1.00 12.92
MOTA	2001	C	ARG	253	13.990	8.732	39.658	1.00 12.93
MOTA	2002	0	ARG	253	12.882	9.268	39.690	1.00 13.28
MOTA	2003	N	ILE	254	15.032	9.268	39.034	1.00 12.41
ATOM	2004		ILE	254	14.927	10.544	38.329	1.00 12.24
MOTA	2005		ILE		16.349	11.025	37.858	1.00 12.27
MOTA	2006			254	1 6.870	10.133	36.704	1.00 12.14
ATOM -	2007			254	16.295	12.496	37.429	1.00 13.31
MOTA	2008	CD1	ILE	254	17.706	13.120	37.107	1.00:12.84
ATOM	2009	С.	ILE	254	13.951	10.432	37.157	1.00 13.19
MOTA	2010	0	ILE	254	13.853	9.384	36.510	1.00 12.84
MOTA	2011	N	THR	255	13.209	11.510	36.909	1.00 12.98
MOTA	2012	CA	THR	255	12.264	11.570	35.804	1.00 14.49
MOTA	2013	CB	THR	255	11.020	12.385	36.186	1.00 15.98
MOTA	2014	0G1	THR	255	11.419	13.721	36.526	1.00 16.76
MOTA	2015	CG2	THR	255	10.342	11.754	37.39 0	1.00 16.72

FIG.11B-48

ATOM	2016	c -	THR	255		12.962	12.266	34.662	1.00 14.89
ATOM	2017		THR	255		14.002	12.908	34.850	1.00 14.36
MOTA	2018	N	ILE	256		12.387	12.180	33.473	1.00 14.96
MOTA	2019	CA	ILE	256		13.022	12.822	32.338	1.00 15.85
ATOM	2020	CB	ILE :	256		12.323	12.446	31.031	1.00 15.90
ATOM		CG2				12.969	13.227	29.886	1.00 16.70
MOTA	2022		•	256	7.7	12.416	10.929	30.824	1.00 16.68
ATOM	2023				•	11.763	10.410	29.490	1.00 17.39
MOTA			ILE					32.495	1.00 15.76
MOTA			ILE	-	11	14.146	14.936		1.00 15.74
ATOM	2026		PRO	257		12.058	15.008	32.988	1.00 16.02
ATOM	2027		PRO	257		10.663	14.579	33.216	1.00 16.52
MOTA	2028		PRO	257		12.196	16.461	33.137	1.00 16.77
ATOM			PRO	257		10.845	16.869	33.730	1.00 16.80
MOTA	2030	CG	PRO	257		9.886	15.903	33.072	1.00 16.27
ATOM	2031		PRO	257		13.448	16.832	34.008	1.00 17.42
ATOM			PRO	257		14.093	17.848	33.765	1.00 18.68
MOTA	2033	N	ASP	258		13.776	15.999	34.996	1.00 17.57
ATOM	2034	CA	ASP	258		14.934	16.252	35.857	1.00 17.40
ATOM	2035	CB	ASP	258		14.727	15.585	37.229	1.00 18.84
MOTA	2036	CG	ASP	258		13.770	16.499	38.040	1.00 20.18
MOTA	2037	OD1	ASP	258		13.098	16.010	38.969	1.00 19.96
ATOM	2038	0D2	ASP			13.710	17.711	37.743	1.00 22.03
ATOM	2039	C	ASP	258		16.254	15.810	35.165	1.00 16.71
ATOM	2040	0	ASP	258		17.313	16.400	35.402	1.00 17.53
MOTA	2041	N	ILE	259	May.	16.180	14.792	34.312	1.00 16.00
MOTA	2042	CA	ILE	259		17.361	14.357	33.567	1.00 14.73
MOTA	2043	CB	ILE	259		17.061	13.134	32.658	1.00 13.25
MOTA	2044	CG2	ILE	259		18.186	12.932	31.636	1.00 11.56
ATOM	2045	CG1	ILE	259	. ≒.	16.926	11.870	33.512	1.00 13.50
MOTA	2046	CD1	ILE:	259	+ 2 + h = -	16.359	10.635	32.718	1.00 14.05
MOTA	2047					17.777	15.511	32.650	1.00 15.65
ATOM	2048	0	ILE	259		18.956	15.768	32.455	1.00 15.76
ATOM	2049	N	LYS	260	-	16.801	16.225	32.097	1.00 17.80
ATOM	2050	CA	LYS	260		17.134	17.318	31.195	1.00 19.60
ATOM	2051	CB	LYS	260		15.882	17.772	30.437	1.00 21.62
ATOM	2052	CG	LYS	260	1	14.787	18.292	31.278	1.00 25.24
MOTA	2053	CD	LYS	260		13.557	18.718	30.460	1.00 27.14
MOTA	2054	CE	LYS	260		12.448	19.266	31.394	1.00 28.28
MOTA	2055	NZ	LYS	260		13.018	20.136	32.457	1.00 28.09
MOTA	2056		LYS	260	ł	17.787	18.495	31.910	1.00 19.77
ATOM	2057			- 260	}	18.302	19.412	31.258	1.00 19.52

FIG.11B-49

MOTA	2058	N	LYS	261	• • •	17.769	18.465	33.244	1.00 19.58
ATOM	2059	CA	LYS.	261	٠.	18.377	19.513	34.063	1.00 20.76
MOTA	2060	CB	LYS	261		17.441	19.911	35.207	1.00 22.07
ATOM	2061	CG	LYS	261		16.225	20.617	34.640	1.00 24.13
MOTA	2062	CD	LYS	261		15.304	20.904	35.853	1.00 25.66
MOTA	2063	CE	LYS	261		13.996	21.718	35.627	1.00 27.76
MOTA	2064	NZ	LYS	261		14.253	23.180	35.441	1.00 29.98
ATOM	2065	C	LYS	261		19.708	19.078	34.677	1.00 19.39
ATOM	2066	0	LYS	261		20.398	19.877	35.320	1.00 20.44
MOTA	2067	N	ASP	262		20.075	17.817	34.461	1.00 17.11
MOTA	2068	CA	ASP	262		21.307	17.258	35.002	1.00 16.68
MOTA	2069	CB	ASP	262		21.348	15.747	34.725	1.00 15.81
ATOM	2070	CG	ASP	262		22.727.	15.133	34.962	1.00 15.09
MOTA	2071	OD1	ASP	262		23.534	15.000	34.021	1.00 15.37
ATOM	2072	OD2	ASP	262		23.049	14.765	36.105	1.00 15.37
MOTA	2073	C	ASP	262		22.539	17.953	34.484	1.00 16.67
MOTA	2074	. 0	ASP	262		22.595	18.357	33.322	1.00 15.94
ATOM	2075	N	ARG	263		23.535	18.094	35.353	1.00 16.49
MOTA	2076	CA	ARG	263		24.781	18.773	34.997	1.00 17.66
MOTA	2077	CB	ARG	263		25.751	18.741	36.179	1.00 19.91
MOTA	2078	CG	ARG	263		27.046	19.552	35.881	1.00 23.77
MOTA	2079	CD	ARG	263	•	27.952	19.734	37.161	1.00 26.98
ATOM	2080	NE	ARG	263		28.878	18.625	37.404	1.00 30.04
MOTA	2081		ARG.	263	· ; :.	28.535	17.410	37.833	
MOTA	2082	NH1	ARG	263		27.264	17.108	38.076	1.00 32.73
MOTA	2083	NH2	ARG	263		29.476	16.495	38.044	1.00 32.79
MOTA	2084	C	ARG	263		25.481	18.182	33.763	1.00 16.93
MOTA	2085	0 .	A RG	263		25.873	18.915	32.858	1.00 17.03
MOTA	2086	N	TRP	264		25.643	16.864	33.725	1.00 15.70
MOTA	2087	CA	TRP	264		26.297	16.256	32.577	1.00 14.73
MOTA	2088	CB	TRP	264		26.576	14.770	32.818	1.00 14.69
MOTA	2089	CG	TRP	264		27.266	14.159	31.637	1.00 13.54
MOTA	2090	CD2	TRP	264		26.677	13.327	30.637	1.00 13.29
MOTA	2091	CE2	TRP	264		27.683	13.043	29.683	•
MOTA	2092	CE3	TRP	264		25.390		30.448	
MOTA	2093	CD1	TRP	264		28.568	14.345		1.00 13.98
ATOM	2094	NE1	TRP	264		28.823		-	1.00 12.82
MOTA	2095	CZ2	TRP	-		27.448		28.556	–
MOTA	2096	CZ3	TRP	264		25.157	11.997	29.329	1.00 11.93
MOTA	2097	CH2	TRP	264		26.178	11.731	28.397	1.00 11.95
ATOM	2098	C	TRP	264		25.437	16.398	31.287	1.00 14.29
MOTA	2099	0	TRP	264		25.954	16.683		1.00 13.06

	••					•			•
MOTA	2100	N TYR	265	2	4.132	16.217	31.427	1.00 1	4.18
MOTA	2101	CA TYR .	265	2	3.228	16.327	30.287	1.00 1	L4.24
MOTA	2102	CB	265	. 2	1.779	16.158	30.753	1.00 1	L4 . 67
MOTA	2103	CG TYR	265	2	0.786	16.027	29.623	1.00 1	14.92
ATOM	2104	CD1 TYR	265	2	0.225	17.150	29.014	1.00 1	L5.83
MOTA	2105	CE1 TYR	265	1	9.303	17.016	27.971	1.00 1	16.00
ATOM	2106	CD2 TYR	265	2	0.404	14.768	29.161	1.00 1	15.34
MOTA	2107	CE2 TYR	265	1	9.492	14.627	28.124	1.00	15.43
ATOM	2108	CZ TYR	265	1	8.948	15.747	27.540	1.00	16.30
ATOM	2109	OH TYR	265	1	8.056	15.596	26.517	1.00	17.21
ATOM	2110	C TYR	265	2	23.420	17.698	29.574	1.00	14.08
ATOM	2111	O TYR	265	2	23.402	17.782	28.340	1.00	13.66
ATOM	2112	N ASN	266	2	237638	18.7 4 3	30.367	1.00	14.50
ATOM	2113	CA ASN	266	2	23.816	20.093	29.832	1.00	15.04
MOTA	2114	CB ASN	266		23.127		30.744	1.00	16.27
ATOM	2115	CG ASN	266		21.623	20.873	30.628	1.00	17.50
ATOM	2116	OD1 ASN	266		21.019	21.164	29.595	1.00	18.67
MOTA		ND2 ASN	266		21.017	20.324	31.689	1.00	17.20
ATOM	2118	C ASN	266		25.283	20.545	29.665	1.00	15.81
ATOM	2119	O ASN	266		25.551	21.696	29.333	1.00	15.28
ATOM	2120	N LYS	267		26.229	19.639	29.867	1.00	15.30
MOTA	2121	CA LYS	267		27.626	20.022	29.739	1.00	16.63
ATOM	2122				28.510	19.009	30.468	1.00	18.26
ATOM	2123	CG LYS	and the second second		29.969	19.316	30.381	1.00	18.95
ATOM	2124	CD LYS			30.607		31.212	1.00	20.54
MOTA	2125	CE LYS			32.097	18.519	31.285	1.00	21.22
ATOM	2126	NZ LYS			32.271	19.896	31.837	1.00	25.63
ATOM	2127	C LYS		rand gr	28.096	20.128	28.280	1.00	17.13
ATOM	2128	0 LYS	267		27.925	19.200	27.491	1.00	17.02
ATOM	2129	N PRO	268	ya T.	28.668	21.285	27.899	1.00	17.19
ATOM	2130		268		28.680	22.567	28.624	1.00	18.04
MOTA	2131				29.151	21.449	26.525	1.00	18.03
MOTA	2132		268		29.594	22.914	26.489	1.00	18.73
MOTA	2133	CG PRO	-				27.485	1.00	18.08
ATOM	2134		268		30.291	20.455	26.275	1.00	19.66
MOTA	2135	O PRO	268		31.263	20.407	27.041	1.00	18.33
MOTA	2136	N LEU	269		30.183	19.688	25.196	1.00	20.60
MOTA	2137		269	. • •	31.191	18.680	24.884	1.00	23.11
ATOM	2138		269		30.735	17.287	25.326	1.00	21.93
ATOM	2139		269		30.429	16.951	26.782	1.00	21.11
ATOM	2140				29.824	15.559	26.864	1.00	20.44
ATOM	2141	CD2 LEU	269		31.709	17.029	27.611	1.00	21.85

FIG.11B-51

•					•			
MOTA	2142	C	LEU	269	31.519	18.481	23.398	1.00 25.42
MOTA	2143	0	LEU	269	32.694	18.357	23.024	1.00 25.84
MOTA	2144	N	LYS	270	30.478	18.456	22.572	1.00 28.13
MOTA	2145	CA	LYS	270	30.638	18.172	21.148	1.00 31.59
MOTA	2146	CB	LYS	270	29.792	16.951	20.777	1.00 32.48
MOTA	2147	CG	LYS	270	29.974	16.336	19.385	1.00 34.36
ATOM	2148	CD	LYS	270	29.245	14.976	19.311	1.00 34.97
ATOM	2149	CE	LYS	270	29.342	14.378	17.904	1.00 36.24
MOTA	2150	NZ	LYS	270	28.578	13.107	17.794	1.00 37.61
ATOM	2151	C	LYS	270	30.326	19.247	20.152	1.00 32.71
MOTA	2152	0	LYS	270	29.331	19.965	20.267	1.00 33.61
MOTA	2153	N	LYS	271	31.182	19.353	19.143	1.00 34.01
MOTA	2154	CA	LYS	271	30.984	20.338	18.093	1.00 34.65
ATOM	2155	CB	LYS	271	32.296	20.639	17.364	1.00 34.43
ATOM	2156	CG	LYS	271	33.612	21.016	18.114	1.00 34.00
MOTA	2157	CD	LYS	271	33.425	21.680	19.493	1.00 32.78
MOTA	2158	CE	LYS	271	34.683	22.342	20.087	1.00 32.43
ATOM	2159	NZ	LYS	271	34.911	23.689	19.493	1.00 29.16
ATOM	2160	C	LYS	271	30.002	19.770	17.100	1.00 35.51
ATOM	2161	0	LYS	271	29.873	18.545	16.970	1.00 35.67
ATOM	2162	N	GLY	272	29.304	20.653	16.392	1.00 36.22
ATOM	2163	CA	GLY	272	28.334	20.207	15.412	1.00 37.58
ATOM	2164	C	GLY	272	28.974	19.331	14.357	1.00 38.77
MOTA		0	GLY	272	•	19.085	14.392	
MOTA	2166	N	ALA	273	28.165	18.859	13.415	1.00 39.07
MOTA	2167	CA	ALA	273	28.663	18.004	12.347	1.00 39.69
MOTA	2168	CB.		273	27.559	17.736	11.334	1.00 39.13
MOTA .	2169	C	ALA	273	29.834	18.652	11.667	1.00 40.12
MOTA	2170	0	ALA	273	30.138	19.821	11.907	1.00 40.27
MOTA	2171	N	ALA	274	30.506	17.889	10.811	
MOTA	2172	CA	ALA	274	31.650	18.406	10.075	
MOTA	2173	CB	ALA	274		17.297	9.834	1.00 41.17
MOTA	2174	C	ALA	274	31.149	10 000	8.761	1.00 41.68
MOTA	2175	0	ALA	274	30.049	18.613	8.317	
MOTA	2176	N	ALA:	275	31.947	19.820	8.143	1.00 41.82
ATOM -	2177	CA	ALA	275	31.600	20.428	6.860	1.00 41.71
ATOM	2178	CB	ALA	275	31.811	19.414	5.741	1.00 41.41
MOTA	2179	Ċ	ALA	275	30.158	20.973	6.807	1.00 41.89
MOTA	2180	0	ALA	275	29.423	20.708	5.850	1.00 42.04
MOTA	2181	N	ALA	276	29.767	21.733	7.829	1.00 41.67
ATOM	2182	CA	ALA	. 276	28.425	22.310	7.881	1.00 41.71
MOTA	2183	CB	ALA	276	27.377	21.201	7.836	1.00 40.99

FIG.11B-52

				. (
MOTA	2184	C .	ALA	276		28.226	23.157	9.127	1.00 41.84
MOTA	2185	0	ALA	276	.*	28.106	24.394	9.001	1.00 42.24
MOTA	2186	OT	ALA	276	,. · ·	28.190	22.590	10.239	1.00 42.70
ATOM	2187		WAT	500	٠,	7.427	-2.493	31.016	1.00 12.44
MOTA	2188	0H2	WAT	501	٠.	7.228	0.472	30.486	1.00 11.40
ATOM	2189		WAT		٠	8.194	1.752	37.455	1.00 11.41
ATOM	2190		WAT	503	1.17	12.286	-2.112	29.696	1.00 12.42
ATOM	2191			504		12.428	-0.037	27.883	1.00 11.16
ATOM	2192	0H2	WAT	505	·	8.356	10.402	31.031	1.00 13.84
ATOM	2193	0H2	WAT	507	i de l	15.558	-3.663	26.632	1.00 12.28
MOTA	2194	0H2	WAT	508	T : 1	6.988	4.420	40.772	1.00 14.28
MOTA	2195	0H2	WAT	509		11.678	7.753	and the second second	1.00 15.05
MOTA	2196	0H2	WAT	510	₽ ₹	9.743	10.806		1.00 13.52
MOTA	2197		WAT	511		14.137	-4.264		1.00 11.96
MOTA	2198	0H2	WAT	512		12.161	-1.918		1.00 17.92
MOTA	2199	0H2	WAT	513	NG.	23.034	-4.599	32.333	
MOTA	2200	OH2	WAT	514		13.701		31.829	
MOTA	2201	OH2	WAT:	515		7.725			
MOTA	2202	OH2	WAT	516	AMEST TO COLO	the second second	8.123		
ATOM	2203	OH2	WAT	517			-2.193		
MOTA	2204	The second of the second of	WAT.	er for the state of		3.854	A resolution of the first		and the first term of the state
MOTA	2205	17 1 1 2 2 .	WAT			6.585		28.016	
MOTA	2206	OH2	WAT	520	100		6.179		
MOTA	2207	OH2	2 WAT			-2.497			
ATOM	2208		2 WAT			25.696	•		
MOTA	2209	100000	2 WAT			10.006			1.00 17.12
MOTA	2210		2 WAT			18.801			1.00 12.85
MOTA	2211	1	2 WAT			9.859		29.813	
MOTA	2212		2 WAT			23.813		12.469	
ATOM	2213	OH	2 WAT	527		33.619			
ATOM	2214	OH	2 WAT	528	N. 4		-0.860	12.686	1.00 16.92
MOTA				529		5.067		•	1.00 15.55
ATOM				530		16.206		45.578	
ATOM	2217	OH	2 WAT	531				39.468	
MOTA						9.848		13.922	
MOTA				533		8.482		and the second second	1.00 21.35
MOTA	2220		2 WAT			1.955			
MOTA	222		2 WAT			8.004			•
ATOM	2222		2 WAT			9.589			
ATOM	2223		2 WAT			13.208	•		
ATOM	2224		2 WAT			12.245	_		· · · · · · · · · · · · · · · · · · ·
MOTA	222	5 OH	2 WAT	540	j	11.065	-3.376	13.747	1.00 23.19

FIG.11B-53

ATOM	2226	OH2 WAT	542		10.329	-0.437	-17.113	1.00 15.49
MOTA	2227	OH2 WAT	543		34.999	12.972	30.493	1.00 20.46
MOTA	2228	OH2 WAT	544		6.038	-4.021	-15.260	1.00 18.16
MOTA	2229	OH2 WAT	545		2.722	-3.465	20.201	1.00 22.45
ATOM	2230	OH2 WAT	546		23.120	17.680	38.118	1.00 21.95
MOTA	2231	OH2 WAT	547		4.224	12.544	29.399	1.00 22.88
MOTA	2232	OH2 WAT	548	•	27.520	19.070	23.817	1.00 18.56
ATOM	2233	OH2 WAT	549		11.453	0.217	-14.778	1.00 18.21
ATOM	2234	OH2 WAT	550		8.159	8.888	13.504	1.00 22.71
MOTA	2235	OH2 WAT	551		7.518	-1.202	14.804	1.00 19.40
MOTA	2236	OH2 WAT	552		25.729		13.336	1.00 25.24
MOTA	2237	OH2 WAT	553		8.421	2.347	13.686	1.00 18.49
MOTA	2238	OH2 WAT	554~	٠.	32.146	14.746	31.790	1.00 16.58
MOTA	2239	OH2 WAT	555		10.588	15.422	22.583	1.00 20.42
MOTA	2240	OH2 WAT	556		-7.789	5.192	30.091	1.00 21.72
MOTA	2241	OH2 WAT	557		24.235	11.751	41.632	1.00 23,21
MOTA	2242	OH2 WAT	558		13.097	5.532	4.167	1.00 22.65
MOTA	2243	OH2 WAT	561		7.327	8.904	36.362	1.00 19.07
MOTA	2244	OH2 WAT	562		5.298	7.204	36.854	1.00 19.10
MOTA	2245	OH2 WAT	563		17.888	14.061	15.698	1.00 28.05
MOTA	2246	OH2 WAT	564		5.803	10.952	34.891	1.00 25.56
MOTA	2247	OH2 WAT	565		19.385	-8.096	22.747	1.00 27.33
MOTA	2248	OH2 WAT	567		-5.961	9.687	24.986	1.00 28.68
ATOM	2249	OH2 WAT	568		12.502	16.572	24.587	1.00 24.90
MOTA	2250	OH2 WAT	569		4.420	13.953	22.823	1.00 19.89
ATOM.	2251	OH2 WAT	570		6.037	16.089	27.263	1.00 27.33
ATOM	2252	OH2 WAT	571		0.295	-4.830		1.00 22.95
MOTA	2253	OH2 WAT	572		5.126	7.073	43.112	1.00 26.68
ATOM	2254		573		7.925	12.617	34.293	1.00 19.25
MOTA	2255	OH2 WAT	574		2.838	8.548	37.282	1.00 22.58
MOTA	2256	OH2 WAT	575		6.541	6.869	39.585	1.00 20.25
ATOM	2257	OH2 WAT	577		16.348	13.014	13.638	1.00 21.44
ATOM	2258	OH2 WAT	578		20.689	-11.863	31.456	1.00 22.24
ATOM	2259	OH2 WAT	579	7 <u>:</u>	28.216	-3.073	39.433	1.00 23.94
MOTA	2260	OH2 WAT	580		4.817	12.316	31.998	1.00 24.78
MOTA	2261	OH2 WAT	582		2.495	10.173	33.047	1.00 25.54
MOTA	2262	OH2 WAT	584		9.873	-9.843	26.499	1.00 22.35
MOTA	2263	OH2 WAT	585		18.849	6.343	6.565	1.00 23.80
ATOM	2264	OH2 WAT	586		5.936	15.554	24.398	1.00 32.00
MOTA	2265	OH2 WAT	5 87		7.942	15.782		•
ATOM	2266	OH2 WAT	588		6.895	14.265	32.126	1.00 25.39
ATOM	2267	OH2 WAT	589	•	-0.295	-3.712	42.925	

					•			•
MOTA	2268	OH2	WAT	590	-3.936	9.005	35.847	1.00 24.10
MOTA	2269	OH2	WAT	591	18.913	2.038	44.494	1.00 26.21
MOTA	2270	OH2	WAT	592	28.625	-6.540	28.424	1.00 26.01
ATOM	2271	OH2	WAT	593	26.141	-9.992	35.885	1.00 25.72
MOTA	2272	OH2	WAT	594	-4.117	0.747	36.348	1.00 21.02
ATOM	2273	OH2	WAT	595	4.898	-5.492	46.334	1.00 25.89
ATOM	2274	OH2	WAT	596	-1.825	-3.880	35.982	1.00 26.80
MOTA	2275	OH2	WAT	597		-10.153		1.00 29.07
MOTA	2276	OH2	WAT	598	6.074	7.250	12.492	1.00 26.52
MOTA	2277	OH2	TAW	599	14.343		-12.155	
ATOM	2278	OH2	WAT	600			35.592	1.00 31.55
MOTA	2279		WAT	601			· ·	1.00 31.89
ATOM	2280	OH2	WAT			16.433		1.00 27.84
ATOM	2281		WAT	604		and the second of the second o		1.00 25.33
ATOM	2282		WAT	605		14.038		1.00 31.51
ATOM		OH2		606	_	/	28.785	
MOTA	2284		A	607			17.394	and the second s
MOTA	2285		WAT	608			-8.218	
ATOM	2286		WAT	609	25.470		Application of the control of the co	1.00 36.33
ATOM	2287		WAT	610	5.918	-3.633		
MOTA	2288	er i de la companya	WAT	611			33.710	
MOTA	2289		WAT	612			37.176	The first term of the first te
MOTA		OH2		613			38.036	
MOTA	2291		WAT	614	22.538			1.00 25.61
MOTA	2292		TAW		2.677		42.779	
MOTA	2293	1.1	WAT		-1.262		1.007	1.00 41.96
MOTA	2294		WAT	617	14.838		15.686	
MOTA	2295		WAT	618	7.254			
MOTA	2296	*	TAW	619	14.437			1.00 24.67
MOTA	2297		WAT	620	13.993		27.899	1.00 37.44
	2298			621			17.703	
MOTA	2299		WAT		1.225		-11.071	
MOTA	2300	. '		623			41.236	
MOTA	2301			624	•		7.878	
MOTA			WAT	625	6.639		16.521	
MOTA	2303		WAT	626			13.973	
MOTA	2304		WAT	627	3.444		-2.127	
MOTA	2305		2 WAT	628	8.270		16.481	1.00 27.56
MOTA	2306		TAW S	629			41.048	
MOTA	2307		2 WAT	630			21.168	
ATOM	2308		2 WAT	631			-11.485	
MOTA	2309	UH2	2 WAT	632	2.999	1.453	-10.337	1.00 32.88

FIG.11B-55

ATOM	2310	0H2 I		633	-10.039	6.144	37.785	1.00 32.06
ATOM	2311	0H2 1		634	25.680	21.534	32.761	1.00 30.38
MOTA	2312	0H2 I	•	636	1.101	14.667	27.285	1.00 33.90
ATOM	2313	0H2 I		637	4.677	-7.995	15.521	1.00 39.69
MOTA	2314	0H2 1		638	-4.199	10.629	27.487	1.00 25.74
MOTA	2315	OH2 1		639	16.727	16.185	23.380	1.00 24.68
ATOM	2316	0H2		641	4.762	8.324	41.074	1.00 32.42
ATOM	2317	0H2	WAT	642	1.346	-0.850	-3.508	1.00 37.14
MOTA	2318	OH2	WAT	643	6.494	-5.448	-2.382	1.00 29.92
MOTA	2319	0H2 1	WAT	644	-0.637	10.395	17.913	1.00 32.31
MOTA	2320	OH2 1	WAT	645	28.896	-3.506	20.216	1.00 28.05
ATOM	2321	OH2	WAT .	646	13.649	-8.354	22.832	1.00 36.52
ATOM	2322	0H2 1	WAT	647	-4.016	-2.000	41.527	1.00 41.51
ATOM	2323	0H2 1	WAT	648	-3.699	4.194	15.863	1.00 34.38
MOTA	2324	OH2	WAT	649	18.236	9.536	44.036	1.00 40.10
MOTA	2325	OH2	WAT	650	-2.251	-2.420	29.819	1.00 37.50
MOTA	2326	OH2	WAT	651	28.245	9.734	16.414	1.00 31.59
MOTA	2327	OH2	WAT	652	25.887	14.410	11.861	1.00 39.37
MOTA	2328	OH2	WAT	653	-4.668	-3.492	21.738	1.00 38.13
ATOM	2329	OH2	WAT	654	15.932	8.831	-4.665	1.00 42.38
MOTA	2330	OH2	WAT	655	39.349	-0.041	40.457	1.00 36.11
MOTA	2331	OH2	WAT	656	16.291	15.362	18.684	1.00 28.74
ATOM	2332	OH2	WAT	657	20.650	8.704	43.546	1.00 26.18
MOTA	2333	OH2	WAT	658	21.731	4.870	-9.446	1.00 41.19
MOTA	2334	OH2	WAT	659	27.579	-8.698	29.528	1.00 36.99
MOTA	2335	OH2	WAT	660	15.065	1.058	-9.945	1.00 34.45
MOTA	2336	C1	ADPN	800	15.589	-7.036	12.366	1.00 29.43
MOTA	2337	C2	ADPN	800	16.795	-6.562	11.567	1.00 27.99
MOTA	2338	01	ADPN	800	16.276	-5.540	10.684	1.00 26.79
MOTA	2339	C3	ADPN	800	17.832	-5.869	12.464	1.00 28.23
MOTA	2340	02	adpn	800	19.138	-6.070	11.920	1.00 29.48
MOTA	2341	C4	adpn	800	17.379	-4.406	12.439	1.00 27.06
MOTA	2342	03	ADPN	800	18.452	-3.550	12.841	1.00 29.12
MOTA	2343	C5	ADPN	800	16.915	-4.275	10.979	1.00 25.84
MOTA	2344	N1	ADPN	800	15.939	-3.162		1.00 23.38
MOTA	2345	C6	ADPN	800	14.655	-3.121	11.351	1.00 23.14
MOTA	2346	N2	ADPN	800	14.038	-1.962	10.976	
MOTA	2347	C7	ADPN	800	14.938	-1.266	10.236	1.00 22.16
MOTA	2348	C8	ADPN	800	14.895	-0.025	9.594	1.00 22.13
ATOM	2349	N3	ADPN	800	13.812	0.764	9.707	1.00 21.32
MOTA	2350		ADPN	800	16.025	0.390	8.889	1.00 21.64
ATOM	2351		ADPN	800	17.152	-0.341	8.819	1.00 21.90
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MOTA	2352	N5	ADPN	800	17.271	-1.548	9.430	1.00 21.98
ATOM	2353	C10	ADPN	800	16.144	-2.011	10.140	1.00 22.86
ATOM	2354	S	S04	901	-0.220	-4.850	27.961	1.00 26.12
ATOM	2355	01	S04	901	0.507	-5.374	26.794	1.00 26.13
ATOM	2356	02	S04	901	0.700	-4.720	29.109	1.00 28.87
ATOM	2357	03	S04	901	-1.308	-5.781	28.341	1.00 24.66
ATOM	2358	04	S04	901	-0.818	-3.538	27.657	1.00 29.71
	•							

FIG.11B-57





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- (71) Applicant: Agouron Pharmaceuticals, Inc. La Jolla, CA 92037 (US)
- (72) Inventors:
 - Chen, PingSan Diego, California 92129 (US)
 - Kan, Chen-Chen, Keck Graduate Inst. of A.L.S.
 Claremont, California 91711 (US)
 - Luo, Chun Irvine, California 92206 (US)
 - Margosiak, Stephen Escondido, California 92025 (US)
 - O'Connor, Patrick
 San Diego, California 92130 (US)

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- Tempczyk-Russel, Anna San Diego, California 92130 (US)
- Nguyen, Binh
 San Diego, California 92130 (US)
- Sarup, Jay Chand
 San Diego, California 92122 (US)
- Gaur, Smita
 San Diego, California 92129 (US)
- Anderson, Mark Brian
 Orinda, California 94563 (US)
- Deng, Ya-Li
 San Diego, California 92130 (US)
- Lundgren, Karen
 San Diego, California 92109 (US)
- Register, James
 San Diego, California 92192 (US)
- (74) Representative: Hofmann, Harald et al Sonnenberg Fortmann, Patent- und Rechtsanwälte, Herzogspitalstrasse 10a 80331 München (DE)
- (54) Catalytic domain of the human effector cell cycle checkpoint protein kinase, Chk1, materials and methods for identification of inhibitors thereof

(57) The present invention relates to the identification, isolation and purification of the catalytic domain of the human effector checkpoint protein kinase (hChk1). A 1.70 crystal structure of the hChk1 kinase domain in the active conformation is reported herein. The kinase domain of hChk1 and its associated crystal structure is described for use in the discovery, identification and

characterization of inhibitors of hChk1. This structure provides a three-dimensional description of the binding site of the hChk1 for structure-based design of small molecule inhibitors thereof as therapeutic agents. Inhibitors of hChk1 find utility in the treatment of hyperproliferative disorders such as HIV and cancer.



EUROPEAN SEARCH REPORT

Application Number EP 00 12 3738

Category	Citation of document with indicati of relevant passages	on, where appropriate,	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int.Cl.7)
X Y	WO 99 11795 A (CARR AND CORP (US)) 11 March 199 * the whole document * Seq. Id. Nos. 1 and 2 s identity to Seq. d. Nos respectively	9 (1999-03-11) show 100% and 99%	1,2, 14-40 3-13,41	C12N15/54 C12N9/12 C12Q1/34
	* claims 1-29; examples	1,6 *		
Х	SANCHEZ ET AL: "Consercheckpoint pathway in m DNA damage to Cdk regul Cdc25"	ammals: linkage of	1,2, 14-40	
	SCIENCE, AMERICAN ASSOC ADVANCEMENT OF SCIENCE, vol. 277, 5 September 1 pages 1497-1501, XP0021	, US, 997 (1997-09-05),		
Y	ISSN: 0036-8075 * the whole document * hChk1 exhibits 100% and Seq. Id. Nos. 1 and 2,	99% identity to	3-13,41	
	* figures 1,4 *			TECHNICAL FIELDS SEARCHED (Int.CI.7)
Y	JOHNSON L N ET AL: "Th for substrate recogniti protein kinases" FEBS LETTERS, ELSEVIER AMSTERDAM, NL, vol. 430, no. 1-2,	on and control by SCIENCE PUBLISHERS,	3-13,41	C12N C12Q
	23 June 1998 (1998-06-2 XP004259128 ISSN: 0014-5793 * table 1 *	3), pages 1-11,		
	* the whole document *			
		-/		
.				
	The present search report has been di			
	Place of search MUNICH	Date of completion of the search 25 July 2002	Petr	Examiner
X : partio Y : partio docu	ATEGORY OF CITED DOCUMENTS cutarly relevant if taken alone cutarly relevant if combined with another ment of the same category nological background	T: theory or principl E: earlier patent doc after the filing dat D: document clied i L: document atted fe	e underlying the in current, but publis le in the application	vention



EUROPEAN SEARCH REPORT

Application Number EP 00 12 3738

Category	Citation of document with Income	dication, where appropriate,	Relevant to claim	CLASSIFICATION OF THE APPLICATION (int.Cl.7)
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	THE AMERICAN ASSOCIATES ARESEARCH. PHILADELPH 1999 PROCEEDINGS OF	HIA, PA, APRIL 10 - 1 F THE ANNUAL MEETING	4,	
	THE AMERICAN ASSOCIA	ATION FOR CANCER		
	XP000891503 * abstract *			
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	CURRENT OPINION IN CURRENT BIOLOGY LTD vol. 6. no. 5. Octo	STRUCTURAL BIOLOGY, ., LONDON, GB, ber 1996 (1996–10),		
	pages 595-603, XP00 ISSN: 0959-440X	2128062		TECHNICAL FIELDS SEARCHED (Int.Cl.7)
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· · ·	Total Milote			
	The present search report has	been drawn up for all claims		
	Place of search	Date of completion of the searc		Examiner
	MUNICH	25 July 2002	Pet	ri, B
X · no	CATEGORY OF CITED DOCUMENTS ritioularly relevant if taken alone riticularly relevant if combined with anc	· E : earlier pate after the filling	Inciple underlying the nt document, but publing date cited in the application	ished on, or

ANNEX TO THE EUROPEAN SEARCH REPORT ON EUROPEAN PATENT APPLICATION NO.

EP 00 12 3738

This annex lists the patent family members relating to the patent documents cited in the above-mentioned European search report. The members are as contained in the European Patent Office EDP file on The European Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

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